430 Rec'd PCT/PTO

International Application No. PCT/AU97/00874

Attorney Docket No.

CULLN23.001APC

9/331631

Page 1

TRANSMITTAL LETTER TO THE UNITED STATES DESIGNATED/ELECTED OFFICE (DO/EO/US) CONCERNING A FILING UNDER 35 USC 371

International Application No.:

Date: June 21, 1990 666 1 7

PCT/AU97/00874

International Filing Date:

December 22, 1997

Priority Date Claimed:

U.S. Application No

December 20, 1996

Title of Invention:

ANTIMICROBIAL PROTEINS

Applicant(s) for DO/EO/US:

John Michael Manners; John Paul Marcus; Kenneth Clifford Goulter;

Jodie Lyn Green; Neil Ivan Bower

Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:

- 1. (X) This is a FIRST submission of items concerning a filing under 35 USC 371.
- 2. () This is a **SECOND** or **SUBSEQUENT** submission of items concerning a filing under 35 USC 371.
- 3. (X) This express request to begin national examination procedures (35 USC 371(f)) at any time rather than delay examination until the expiration of the applicable time limit set in 35 USC 371(b) and PCT Articles 22 and 39(1).
- 4. (X) A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date.
- 5. (X) A copy of the International Application as filed (35 USC 371(c)(2))
  - a) () is transmitted herewith (required only if not transmitted by the International Bureau).
  - b) (X) has been transmitted by the International Bureau.
  - c) is not required, as the application was filed in the United States Receiving Office (RO/US).
- 6. () A translation of the International Application into English (35 USC 371(c)(2)).
- 7. (X) Amendments to the claims of the International Application under PCT Article 19 (35 USC 371(c)(3))
  - a) () are transmitted herewith (required only if not transmitted by the International Bureau).
  - b) () have been transmitted by the International Bureau.
  - c) have not been made; however, the time limit for making such amendments has NOT expired.
  - d) (X) have not been made and will not be made.
- 8. () A translation of the amendments to the claims under PCT Article 19 (35 USC 371(c)(3)).
- 9. (X) An oath or declaration of the inventor(s) (35 USC 371(c)(4)).
- 10. (X) A copy of the International Preliminary Examination Report with any annexes thereto, such as any amendments made under PCT Article 34.
- 11. () A translation of the annexes, such as any amendments made under PCT Article 34, to the International Preliminary Examination Report under PCT Article 36 (35 USC 371(c)(5)).

Date: June 21, 1999 Page 2

## Items 11. to 16. below concern other document(s) or information included:

- 12. () An Information Disclosure Statement under 37 CFR 1.97 and 1.98.
- 13. (X) An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included.
- 14. (X) A FIRST preliminary amendment.
  - () A SECOND or SUBSEQUENT preliminary amendment.
- 15. () A substitute specification.
- 16. () A power of attorney and/or address letter.
- 17. (X) International Application as published.
- 18. () Small Entity Statement.
- 19. (X) Sequence listing pages 34-58.
- 20. (X) A diskette containing sequence listing. I hereby certify that the data on the enclosed disk is identical to the sequence listing in the application filed herewith, as required by 37 C.F.R. §1.821(f).
- 21. (X) PCT request form.
- 22. (X) A return prepaid postcard.
- 23. (X) The following fees are submitted:

				FEES
	BASIC FEE			\$970
CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE	
Total Claims	41 - 20 =	21 ×	\$18	\$378
Independent Claims	4 - 3 =	1 ×	\$78	\$78
Multiple dependent claims(s	) (if applicable)		\$260	\$0
	TOTAL NATIO	ONAL FEE		\$1426
TOTAL FEES ENCLOSED			\$1426	

- 24. (X) A check in the amount of \$1426 to cover the above fees is enclosed.
- 25. (X) Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31). \$40 per property.

## International Application No. PCT/AU97/00874

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Date: June 21, 1999

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26. (X) The Commissioner is hereby authorized to charge only those additional fees which may be required to avoid abandonment of the application, or credit any overpayment to Deposit Account No. 11-1410. A duplicate copy of this sheet is enclosed.

NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137(a) or (b)) must be filed and granted to restore the application to pending status.

SEND ALL CORRESPONDENCE TO:

KNOBBE, MARTENS, OLSON & BEAR, LLP 620 Newport Center Drive Sixteenth Floor Newport Beach, CA 92660 Signature

Daniel E. Altman Printed Name

34,115

Registration Number

E:\DOCS\DEA\DEA-1300.DOC:kc 062199

PATENT

## IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant	:	Manners, J. et al	) Group Art Unit Unknown
Int'l Appl. No.	:	PCT/AU97/00874	) ) 1
Int'l Filing Date	:	December 22, 1997	)
For	:	ANTIMICROBIAL PROTEINS	)
Examiner	:	Unknown	<i>)</i> )

## PRELIMINARY AMENDMENT

Assistant Commissioner for Patents Washington, D.C. 20231

## Dear Sir:

Preliminary to examination on the merits please amend the above-identified patent application as follows:

## **IN THE SPECIFICATION:**

On page 1 of the Specification, after the Title of the Invention ending on line 2 and before the Field of the Invention statement starting on line 3, please insert --This is the U.S. national phase under 35 U.S.C. § 371 of International application PCT/AU97/00874, filed December 22, 1997, which claims priority to Australian application PO 4275, filed December 20, 1996.--.

On page 4, at line 22, after the phrase "C-3X-C" and before the word "wherein", please insert --(SEQ ID NOs:31-33)--.

On page 4, at line 24, after the phrase "2X-C-3X-C-(10-12)X-C-3X-C-3X-Z" and before the word "wherein", please insert --(SEQ ID NOs:34-36)--.

On page 4, at line 26, after the phrase "C-3X-C-(10-12)X-C-3X-C", please insert --(SEQ ID NOs:37-39)--.

On page 11, at line 15, please cancel the word "synthesise" and substitute in its place -- synthesize--.

On page 13, at line 31, please cancel the word "synthesised" and substitute in its place -- synthesized--.

On page 14, at line 3, please cancel the word "collectivelly" and substitute in its place -- collectively--.

On page 14, at line 10, before the word "which" and after the word "motifs", please insert --(SEQ ID NO:40)--.

On page 16, at line 3, before the word "any" and after the word "remove" please delete the word "the".

On page 20, at line 7, before the word "will" and after the word "spacing", please insert -- (SEQ ID NO:40)--.

On page 22, at line 26, after the phrase "XXX-C-(10-12X)-C-XXX-C" and before the word "where", please insert --(SEQ ID NOs:37-39)--.

On page 22, at line 32, before the word "are" and after the word "segments" please delete the words "is the".

On page 23, at line 4, after the phrase "Z-XX-C-XXX-C-(10-12)X-C-XXX-C-XXX-Z" and before the word "wherein", please insert --(SEQ ID NOs:34-36)--.

On page 24 at line 8, after the "3" and before the ";" please insert --(SEQ ID NO:17)--.

On page 24 at line 11, after the "3" and before the "." please insert --(SEQ ID NO:18)--.

On page 24 at line 16, after the "3" and before the ";" please insert --(SEQ ID NO:19)--.

On page 24 at line 20, after the "3" and before the "." please insert --(SEQ ID NO:20)--.

On page 67 please delete the word "CLAIMS" and substitute therefor --WHAT IS CLAIMED IS--.

## IN THE CLAIMS:

## Please amend the following claims:

- 1. (Amended) A protein fragment having antimicrobial activity, wherein said protein fragment is [selected from:
  - (i) a polypeptide having an amino acid sequence selected from:

residues 29 to 73 of SEQ ID NO: 1

residues 74 to 116 of SEQ ID NO: 1

> residues 117 to 185 of SEQ ID NO: 1 residues 186 to 248 of SEQ ID NO: 1 residues 29 to 73 of SEQ ID NO: 3 residues 74 to 116 of SEQ ID NO: 3 residues 117 to 185 of SEQ ID NO: 3 residues 186 to 248 of SEQ ID NO: 3 residues 1 to 32 of SEQ ID NO: 5 residues 33 to 75 of SEQ ID NO: 5 residues 76 to 144 of SEQ ID NO: 5 residues 145 to 210 of SEQ ID NO: 5 residues 34 to 80 of SEQ ID NO: 7 residues 81 to 140 of SEQ ID NO: 7 residues 33 to 79 of SEQ ID NO: 8 residues 80 to 119 of SEQ ID NO: 8 residues 120 to 161 of SEQ ID NO: 8 residues 32 to 91 of SEQ ID NO: 21 residues 25 to 84 of SEQ ID NO: 22 residues 29 to 94 of SEQ ID NO: 23 residues 31 to 85 of SEQ ID NO: 24 residues 1 to 23 of SEQ ID NO: 25 residues 1 to 17 of SEQ ID NO: 26 residues 1 to 28 of SEQ ID NO: 27

- (ii) a homologue of (i);
- (iii)] a polypeptide containing a relative cysteine spacing of C-2X-C-3X-C-(10-12)X-C-3X-C\_(SEQ ID NOs:37-39) wherein X is any amino acid residue, and C is cysteine[;
- (iv) a polypeptide containing a relative cysteine and tyrosine/phenylalanine spacing of Z-2X-C-3X-C-(10-12)X-C-3X-C-3X-Z wherein X is any amino acid residue, and C is cysteine, and Z is tyrosine or phenylalanine;
- (v) a polypeptide containing a relative cysteine spacing of C-3X-C-(10-12)X-C-3X-C wherein X is any amino acid residue, and C is cysteine;

(vi) a polypeptide with substantially the same spacing of positively charged residues relative to the spacing of cysteine residues as (i); and

- (vii) a fragment of the polypeptide of any one of (i) to (vi) which has substantially the same anitmicrobial activity as (i)].
- 2. (Amended) An isolated or purified protein containing at least one polypeptide fragment according to claim 1, wherein said polypeptide fragment has a sequence selected from within a sequence comprising SEQ ID NO: 1, SEQ ID NO: 3, or SEQ ID NO: 5.
- 3. (Amended) An isolated or purified protein having a sequence selected from SEO ID NO:1, SEO ID NO: 3, or SEQ ID NO:5.
- 7. (Amended) A transgenic plant [harbouring]harboring a DNA construct according to claim 6.
- 9. (Amended) The transgenic plant according to claim 7, wherein said plant is selected from the group consisting of; maize, banana, peanut, field peas, sunflower, tomato, canola, tobacco, wheat, barley, oats, potato, soybeans, cotton, carnations, roses, [or]and sorghum.
- 10. (Amended) The [R]reproductive material of [a]the transgenic plant according to claim 7.
- 11. (Amended) A composition comprising an antimicrobial protein according to claim 1 together with an agriculturally-acceptable carrier diluent or excipient.
- 12. (Amended) A composition comprising an antimicrobial protein according to claim 1 together with a pharmaceutically-acceptable carrier diluent or excipient.
- 13. (Amended) A method of controlling microbial infestation of a plant, the method comprising[:
- (i)] treating said plant with an antimicrobial protein according to claim 1 [or a composition according to claim 11; or
  - (ii) introducing a DNA construct according to claim 6 into said plant].
- 14. (Amended) A method of controlling microbial infestation of a mammal[ian animal], said[the] method comprising treating the [animal]mammal with an antimicrobial protein according to claim 1 [or a composition according to claim 12].
- 15. (Amended) The method of claim 14, wherein said mammal[ian animal] is a human.

- 16. (Amended) A method of preparing an antimicrobial protein, [which]said method [comprises the steps of:] comprising;
- a) obtaining or designing an amino acid sequence which forms a helix-turn-helix structure;
- b) replacing individual residues to achieve substantially the same distribution of positively charged residues and cysteine residues [as in one or more of the amino acid sequences shown in figure 4]; and
- c) [synthesising]synthesizing a protein comprising said amino acid sequence chemically or by recombinant DNA techniques in liquid culture]; and
- d) if necessary, forming disulphide linkages between said cysteine residues].

## Please add the following claims:

- 17. The protein fragment of Claim 1, wherein said protein fragment is a polypeptide containing a relative cysteine and tyrosine/phenylalanine spacing of Z-2X-C-3X-C-(10-12)X-C-3X-C-3X-Z (SEQ ID NOs:34-36) wherein X is any amino acid residue, and C is cysteine, and Z is tyrosine or phenylalanine.
- 18. The protein fragment of Claim 1, wherein said protein fragment is a polypeptide containing a relative cysteine spacing of C-2X-C-3X-C-(10-12)X-C-3X-C-2X-C (SEQ ID NOs:31-33) wherein X is any amino acid residue, and C is cysteine.
- 19. The protein fragment of Claim 1, wherein said protein fragment is selected from the group consisting of:

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residues 29 to 73 of SEQ ID NO: 1; residues 74 to 116 of SEQ ID NO: 1; residues 117 to 185 of SEQ ID NO: 1; residues 186 to 248 of SEQ ID NO: 1; residues 29 to 73 of SEQ ID NO: 3; residues 74 to 116 of SEQ ID NO: 3; residues 117 to 185 of SEQ ID NO: 3; residues 186 to 248 of SEQ ID NO: 3; residues 33 to 75 of SEQ ID NO: 5; residues 76 to 144 of SEQ ID NO: 5;
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residues 145 to 210 of SEQ ID NO: 5; residues 34 to 80 of SEQ ID NO: 7; residues 81 to 140 of SEQ ID NO: 7; residues 33 to 79 of SEQ ID NO: 8; residues 80 to 119 of SEQ ID NO: 8; residues 120 to 161 of SEQ ID NO: 8; residues 32 to 91 of SEQ ID NO: 21; residues 25 to 84 of SEQ ID NO: 22; residues 29 to 94 of SEQ ID NO: 23; residues 31 to 85 of SEQ ID NO: 24; and residues 1 to 23 of SEQ ID NO: 25.
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- 20. The protein fragment of Claim 1 which is truncated, but wherein said truncated protein fragment has substantially the same antimicrobial activity as the nontruncated protein fragment.
- 21. A protein fragment according to Claim 1, wherein the protein fragment is a homologue of a protein fragment selected from the group consisting of:

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residues 29 to 73 of SEQ ID NO: 1; residues 74 to 116 of SEQ ID NO: 1; residues 117 to 185 of SEQ ID NO: 1; residues 186 to 248 of SEQ ID NO: 1; residues 29 to 73 of SEQ ID NO: 3; residues 74 to 116 of SEQ ID NO: 3; residues 117 to 185 of SEQ ID NO: 3; residues 117 to 185 of SEQ ID NO: 3; residues 186 to 248 of SEQ ID NO: 3; residues 33 to 75 of SEQ ID NO: 5; residues 76 to 144 of SEQ ID NO: 5; residues 145 to 210 of SEQ ID NO: 5; residues 34 to 80 of SEQ ID NO: 7; residues 81 to 140 of SEQ ID NO: 7; residues 33 to 79 of SEQ ID NO: 8; residues 80 to 119 of SEQ ID NO: 8;
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residues 120 to 161 of SEQ ID NO: 8; residues 32 to 91 of SEQ ID NO: 21; residues 25 to 84 of SEQ ID NO: 22; residues 29 to 94 of SEQ ID NO: 23; residues 31 to 85 of SEQ ID NO: 24; and residues 1 to 23 of SEQ ID NO: 25.

22. A protein fragment having antimicrobial activity, wherein said protein fragment is selected from the group consisting of:

residues 1 to 32 of SEQ ID NO:5; residues 1 to 23 of SEQ ID NO:26; residues 1 to 17 of SEQ ID NO:27; and residues 1 to 28 of SEQ ID NO:28.

- 23. A homologue of any of the protein fragments of Claim 22.
- 24. An isolated or synthetic DNA encoding a polypeptide fragment according to claim 22.
- 25. A DNA construct which includes a DNA according to claim 24 operatively linked to elements for the expression of said encoded protein.
  - 26. A transgenic plant harboring a DNA construct according to claim 25.
- 27. The transgenic plant according to claim 26, wherein said plant is a monocotyledonous plant or a dicotyledonous plant.
- 28. The transgenic plant according to claim 26, wherein said plant is selected from the group consisting of; maize, banana, peanut, field peas, sunflower, tomato, canola, tobacco, wheat, barley, oats, potato, soybeans, cotton, carnations, roses, and sorghum.
  - 29. The reproductive material of the transgenic plant according to claim 26.
- 30. A composition comprising an antimicrobial protein according to claim 22 together with an agriculturally-acceptable carrier diluent or excipient.
- 31. A composition comprising an antimicrobial protein according to claim 22 together with a pharmaceutically-acceptable carrier diluent or excipient.
- 32. A method of controlling microbial infestation of a plant or mammal, said method comprising treating said plant or mammal with an antimicrobial protein according to claim 22.
  - 33. The method of Claim 32 wherein said mammal is a human.

34. A method of controlling microbial infestation of a plant, the method comprising treating said plant with a composition according to claim 11.

35. A method of controlling microbial infestation of a plant, the method comprising introducing a DNA construct according to claim 6 into said plant.

36. A method of controlling microbial infestation of a mammal, the method comprising treating the mammal with a composition according to claim 12.

37. The method of claim 19, wherein said mammal is a human.

38. A method of controlling microbial infestation of a plant, the method comprising introducing a DNA construct according to claim 25 into said plant.

39. A method of controlling microbial infestation of a mammal, the method comprising treating the mammal with a composition according to claim 30.

40. The method of claim 39, wherein said mammal is a human.

41. The method of claim 16, further comprising forming disulphide linkages between said cysteine residues.

## **IN THE SEQUENCE LISTING:**

Please cancel from the application original Sequence Listing pages 34-66 and substitute therefor the attached Replacement Sequence Listing pages 34-58. Please consecutively renumber all pages following the Replacement Sequence Listing.

#### **REMARKS**

This Supplemental Preliminary Amendment conforms the Sequence Listing of the priority International Patent Application to the rules of practice specified by the U.S. Patent and Trademark Office. Enclosed herewith are: (1) a paper copy of the Replacement Sequence Listing, and (2) a computer readable version of the Replacement Sequence Listing.

The Specification has been amended to include a reference to the PCT application PCT/AU97/00874, filed December 22, and the priority application. The Specification and Claims have been amended to conform to the rules of practice as specified by the United States PTO and to correct minor informalities. Claims 1-3, 7, 9-16 have been amended. Claims 17-40 have been added. Therefore, Claims 1-40 remain pending. Additionally, the amendment directs

entry of the paper copy of the Listing into the application. In view of the foregoing, the application is believed to fully comply with the Sequence Listing disclosure requirements.

## VERIFICATION UNDER 37 C.F.R. §1.821(f) & (g)

All of the sequences in the attached Sequence Listing were included in the application as filed. Pursuant to 37 C.F.R. §1.821(g), no new matter is being added herewith. As required under 37 C.F.R. §1.821(f), I hereby verify that the data on the enclosed disk and the paper copies of the Sequence Listing are identical.

## Conclusion

Should there be any questions concerning this application, the Examiner is respectfully invited to contact the undersigned at the telephone number appearing below.

Respectfully submitted,

KNOBBE, MARTENS, OLSON & BEAR, LLP

Dated: 21 June 1999

By: \( \times \text{Aurel} \)
Daniel E. Altman

Registration No. 34,115

Attorney of Record

620 Newport Center Drive

Sixteenth Floor

Newport Beach, CA 92660

(949) 760-0404

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# ANTIMICROBIAL PROTEINS TECHNICAL FIELD

This invention relates to isolated proteins which exert inhibitory activity on the growth of fungi and bacteria, which fungi and bacteria include some microbial pathogens of plants and animals. The invention also relates to recombinant genes which include sequences encoding the proteins, the expression products of which recombinant genes can contribute to plant cells or cells of other organism's defence against invasion by microbial pathogens. The invention further relates to the use of the proteins and/or genes encoding the proteins for the control of microbes in human and veterinary clinical conditions.

#### **BACKGROUND ART**

Microbial diseases of plants are a significant problem to the agricultural and horticultural industries. Plant diseases in general cause millions of tonnes of crop losses annually with fungal and bacterial diseases responsible for significant portions of these losses. One possible way of combating fungal and bacterial diseases is to provide transgenic plants capable of expressing a protein or proteins which in some way increase the resistance of the plant to pathogen attack. A simple strategy is to first identify a protein with antimicrobial activity *in vitro*, to clone or synthesise the DNA sequence encoding the protein, to make a chimaeric gene construct for efficient expression of the protein in plants, to transfer this gene to transgenic plants and to assess the effect of the introduced gene on resistance to microbial pathogens by comparison with control plants.

The first and most important step in the strategy for disease control described above is to identify, characterise and describe a protein with strong antimicrobial activity. In recent years, many different plant proteins with antimicrobial and/or antifungal activity have been identified and described. These proteins have been categorised into several classes according to either their presumed mode of action and/or their amino acid sequence homologies. These classes include the following: chitinases (Roberts, W.K. et al. [1986] Biochim. Biophys. Acta 880:161-170); β-1,3-glucanases (Manners, J.D. et al. [1973] Phytochemistry 12:547-553); thionins (Bolmann, H. et al. [1988] EMBO J. 7:1559-1565 and Fernadez de Caleya, R. et al. [1972] Appl. Microbiol. 23:998-1000); permatins (Roberts, W. K. et al. [1990] J. Gen. Microbiol. 136:1771-1778 and Vigers, A.J. et al. [1991] Mol. Plant-Microbe Interact. 4:315-323); ribosome-inactivating proteins (Roberts, W. K. et al. [1986] Biochim. Biophys. Acta 880:161-170 and Leah, R. et al. [1991] J. Biol. Chem. 266:1564-1573); plant defensins (Terras, F. R. G. et al. [1995] The Plant Cell 7:573-588); chitin binding proteins (De Bolle, M.F.C. et al. [1992] Plant Mol. Biol. 22:1187-1190 and Van Parijs, J. et al. [1991] Planta 183:258-264); thaumatin-like, or osmotin-like proteins (Woloshuk, C.P. et al. [1991] The Plant Cell 3:619-628 and Hejgaard, J. [1991] FEBS Letts. 291:127-131); PR1-type

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proteins (Niderman, T. et al. [1995] Plant Physiol. 108:17-27.) and the non-specific lipid transfer proteins (Terras, F.R.G. et al. [1992] Plant Physiol. 100:1055-1058 and Molina, A. et al. [1993] FEBS Letts. 3166:119-122). Another class of antimicrobial proteins from plants is the knottin or knottin-like antimicrobial proteins (Cammue, B.P.A. et al. [1992] J. Biol. Chem. 67:2228-2233; Broekaert W.F. et al. (1997) Crit. Rev. in Plant Sci. 16(3):297-323). A class of antimicrobial proteins termed 4-cysteine proteins has also been reported in the literature which class includes Maize Basic Protein (MBP-1) (Duvick, J.P. et al. [1992] J. Biol. Chem. 267:18114-18120). A novel antimicrobial protein which does not fit into any previously described class of antimicrobial proteins has also been isolated from the seeds of Macadamia integrifolia termed MiAMP1 (Marcus, J.P. et al. [1997] Eur. J. Biochem. 244:743-749). In addition, plants are not the sole source of antimicrobial proteins and there are many reports of the isolation of antimicrobial proteins from animal and microbial cells (reviewed in Gabay, J.E. [1994] Science 264:373-374 and in "Antimicrobial peptides"

There is evidence that the ectopic expression of genes encoding proteins that have *in vitro* antimicrobial activity in transgenic plants can result in increased resistance to microbial pathogens. Examples of this engineered resistance include transgenic plants expressing genes encoding: a plant chitinase, either alone (Broglie, K. *et al.* [1991] *Science* 254:1194-1197) or in combination with a β-1,3-glucanase (Van den Elzen, P.J.M. *et al.* [1993] *Phil. Trans. Roy. Soc.* 342:271-278); a plant defensin (Terras, F.R.G. *et al.* [1995] *The Plant Cell* 7:573-588); an osmotin-like protein (Liu, D. *et al.* [1994] *Proc. Natl. Acad. Sci. USA* 91:1888-1892); a PR1-class protein (Alexander, D. *et al.* [1993] *Proc. Natl. Acad. Sci. USA* 90:7327-7331) and a ribosome-inactivating protein (Logemann, J. *et al.* [1992] *Bio/Technology* 10:305-308).

[1994] CIBA Foundation Symposium 186, John Wiley and Sons Publ., Chichester, UK).

Although the potential use of antimicrobial proteins for engineering disease resistance in transgenic plants has been described extensively, there are other applications which are worthy of mention. Firstly, highly potent antimicrobial proteins can be used for the control of plant disease by direct application (De Bolle, M.F.C. et al. [1993] in *Mechanisms of Plant Defence Responses*, B. Fritig and M. Legrand eds., Kluwer Acad. Publ., Dordrecht, NL, pp. 433-436). In addition, antimicrobial peptides have potential therapeutic applications in human and veterinary medicine. Although this has not been described for peptides of plant origin it is being actively explored with peptides from animals and has reached clinical trials (Jacob, L. and Zasloff, M. [1994] in "Antimicrobial Peptides", *CIBA Foundation Symposium 186*, John Wiley and Sons Publ., Chichester, UK, pp. 197-223).

Antimicrobial proteins exhibit a variety of three-dimensional structures which will determine in large part the activity which they manifest. Many of the global structures exhibited by these

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proteins have been determined (Broekaert W.F. et al. (1997) Crit. Rev. in Plant Sci. 16(3):297-323). A large factor in determining the stability of these proteins is the presence of disulfide bridges between various cysteines located in  $\alpha$ -helical and  $\beta$ -sheet regions. Many peptides with toxic activity such as conotoxin are well known to be stabilized by disulfide bridges (see for example Hill, J.M. et al. (1996) Biochemistry 35(27): 8824-8835). In the case of the conotoxin referenced above, a compact structure is formed consisting of a helix, a small -hairpin, a cis-hydroxyproline, and several turns. The molecule is stabilized by three disulfide bonds, two of which connect the  $\alpha$ -helix and the  $\beta$ -sheet, forming a solid structural core. Interestingly, eight arginine and lysine side chains in this molecule project into the solvent in a radial orientation relative to the core of the molecule. These cationic side chains form potential sites of interaction with anionic sites on pathogen membranes (Hill, J.M. et al. supra).

The invention described herein constitutes previously undiscovered and thus novel proteins with antimicrobial activity. These proteins can be isolated from *Macadamia integrifolia* (Mi) seeds or from cotton or cocoa seeds. In addition, protein fragments which are antifungal can be derived from larger seed storage proteins containing regions of substantial similarity to the antimicrobial proteins from macadamia described here. Examples of seed storage proteins which contain regions similar to the proteins which have been purified can be seen in Figure 4. *Macadamia integrifolia* belongs to the family Proteaceae. *M. integrifolia*, also known as Bauple Nut or Queensland Nut, is considered by some to be the world's best edible nut. Cotton (*Gossypium hirsutum*) belongs to the family Malvaceae and is cultivated extensively for its fiber. Cocoa (*Threobroma cacao*) belongs to the family Sterculiaceae and is used around the world for a wide variety of cocoa products.

The fact that both the macadamia and cocoa antimicrobial proteins are found in edible portions of these plants makes these peptides attractive for use in genetic engineering for disease resistance since trangenic plants expressing these proteins are unlikely to show added toxicity. Proteins may also be safe for human and veterinary use.

#### SUMMARY OF THE INVENTION

According to a first embodiment of the invention, there is provided a protein fragment having antimicrobial activity, wherein said protein fragment is selected from:

(i) a polypeptide having an amino acid sequence selected from:

residues 29 to 73 of SEQ ID NO: 1

residues 74 to 116 of SEQ ID NO: 1

residues 117 to 185 of SEQ ID NO: 1

residues 186 to 248 of SEQ ID NO: 1

residues 29 to 73 of SEQ ID NO: 3

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			4
			residues 74 to 116 of SEQ ID NO: 3
			residues 117 to 185 of SEQ ID NO: 3
			residues 186 to 248 of SEQ ID NO: 3
			residues 1 to 32 of SEQ ID NO: 5
•	5		residues 33 to 75 of SEQ ID NO: 5
			residues 76 to 144 of SEQ ID NO: 5
<b>A</b>			residues 145 to 210 of SEQ ID NO: 5
			residues 34 to 80 of SEQ ID NO: 7
			residues 81 to 140 of SEQ ID NO: 7
	10		residues 33 to 79 of SEQ ID NO: 8
			residues 80 to 119 of SEQ ID NO: 8
; s = 1;			residues 120 to 161 of SEQ ID NO: 8
			residues 32 to 91 of SEQ ID NO: 21
			residues 25 to 84 of SEQ ID NO: 22
	15		residues 29 to 94 of SEQ ID NO: 24
			residues 31 to 85 of SEQ ID NO: 25
: 144			residues 1 to 23 of SEQ ID NO: 26
			residues 1 to 17 of SEQ ID NO: 27
M			residues 1 to 28 of SEQ ID NO: 28;
	20	(ii)	a homologue of (i);
		(iii)	a polypeptide containing a relative cysteine spacing of C-2X-C-3X-C-(10-12)X-C-3X-
			C-3X-C wherein X is any amino acid residue, and C is cysteine;
		(iv)	a polypeptide containing a relative cysteine and tyrosine/phenylalanine spacing of Z-
			2X-C-3X-C-(10-12)X-C-3X-C-3X-Z wherein X is any amino acid residue, and C is
	25		cysteine, and Z is tyrosine or phenylalanine;
		(v)	a polypeptide containing a relative cysteine spacing of C-3X-C-(10-12)X-C-3X-C
			wherein X is any amino acid residue, and C is cysteine;
		(vi)	a polypeptide with substantially the same spacing of positively charged residues
			relative to the spacing of cysteine residues as (i); and
	30	(vii)	a fragment of the polypeptide of any one of (i) to (vi) which has substantially the same
9			antimicrobial activity as (i).
		Accor	ding to a second embodiment of the invention, there is provided a protein containing at

According to a second embodiment of the invention, there is provided a protein containing at least one polypeptide fragment according to the first embodiment, wherein said polypeptide fragment

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has a sequence selected from within a sequence comprising SEQ ID NO: 1, SEQ ID NO: 3 or SEQ ID NO: 5.

According to a third embodiment of the invention, there is provided a protein having a sequence selected from SEQ ID NO: 1, SEQ ID NO: 3 or SEQ ID NO: 5.

According to a fourth embodiment of the invention, there is provided an isolated or synthetic DNA encoding a protein according to the first embodiment

According to a fifth embodiment of the invention, there is provided a DNA construct which includes a DNA according to the fourth embodiment operatively linked to elements for the expression of said encoded protein.

According to a sixth embodiment of the invention, there is provided a transgenic plant harbouring a DNA construct according to the fifth embodiment.

According to a seventh embodiment of the invention, there is provided reproductive material of a transgenic plant according to the sixth embodiment.

According to an eighth embodiment of the invention, there is provided a composition comprising an antimicrobial protein according to the first embodiment together with an agriculturally-acceptable carrier diluent or excipient.

According to a ninth embodiment of the invention, there is provided a composition comprising an antimicrobial protein according to the first embodiment together with an pharmaceuticallyacceptable carrier diluent or excipient.

According to a tenth embodiment of the invention, there is provided a method of controlling microbial infestation of a plant, the method comprising:

- treating said plant with an antimicrobial protein according to the first embodiment or a i) composition according to the eighth embodiment; or
- introducing a DNA construct according to the fifth embodiment into said plant. ii)

According to an eleventh embodiment of the invention, there is provided a method of controlling microbial infestation of a mammalian animal, the method comprising treating the animal with an antimicrobial protein according to the first embodiment or a composition according to the ninth embodiment.

According to a twelfth embodiment of the invention, there is provided a method of preparing an antimicrobial protein, which method comprises the steps of:

- obtaining or designing an amino acid sequence which forms a helix-turn-helix structure; a)
- replacing individual residues to achieve substantially the same distribution of positively b) charged residues and cysteine residues as in one or more of the amino acid sequences shown in Figure 4;

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synthesising a protein comprising said amino acid sequence chemically or by

c) recombinant DNA techniques in liquid culture; and

if necessary, forming disulphide linkages between said cysteine residues. d)

Other embodiments of the invention include methods for producing antimicrobial protein.

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#### BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows the results of cation-exchange chromatography of the basic protein fraction of a Macadamia integrifolia extract with the results of a bioassay for antimicrobial activity shown for fractions in the region of MiAMP2c elution.

Figure 2 shows the results of including 1 mM Ca<sup>2+</sup> in a parallel bioassay of fractions from the cation-exchange separation.

Figure 3 shows a reverse-phase HPLC profile of highly inhibitory fractions containing MiAMP2c from the cation-exchange separation in Figure 1 and 2 together with % growth inhibition exhibited by the HPLC fractions.

Figure 4 shows the amino acid sequences of MiAMP2a, b, c and d and protein fragments derived from other seed storage proteins which contain regions of homology to the MiAMP2 series of antimicrobial proteins.

Figure 5 shows an example of a synthetic nucleotide sequence which can be used for the expression and secretion of MiAMP2c in transgenic plants.

Figure 6 shows the alignment of clones 1-3 from macadamia containing MiAMP2a, b, c and d subunits together with sequences from cocoa and cotton vicilin seed storage proteins which exhibit significant homology to the macadamia clones.

Figure 7 displays a series of secondary structure predictions for MiAMP2c.

Figure 8 shows a three-dimensional model of the MiAMP2c protein.

Figure 9 shows stained SDS-PAGE gels of protein fractions at various stages in the expression and purification of TcAMP1 (Theobroma cacao subunit 1), MiAMP2a, MiAMP2b, MiAMP2c and MiAMP2d expressed in E.coli liquid culture.

Figure 10 shows the reverse-phase HPLC purification of cocoa subunit 2 (TcAMP2) after the initial purification step using Ni-NTA media.

Figure 11 shows a western blot of crude protein extracts from various plant species using rabbit antiserum raised to MiAMP2c.

Figure 12 shows a cation-exchange fractionation of the Stenocarpus sinuatus basic protein fraction along with the accompanying western blot which shows the presence of immunologicallyrelated proteins in a range of fractions.

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Figure 13 shows a reverse-phase HPLC separation of *Stenocarpus sinuatus* cation-exchange fractions which had previously reacted with MiAMP2c antibodies (see Figure 14). A western blot is also presented which reveals the presence of putative MiAMP2c homologues in individual HPLC fractions.

Figure 14 is a map of the binary vector pPCV91-MiAMP2c as an example of a vector that can be used to express these antimicrobial proteins in transgenic plants.

Figure 15 shows a western blot to detect MiAMP2c expressed in transgenic tobacco plants.

## BEST MODE AND OTHER MODES FOR CARRYING OUT THE INVENTION

The following abbreviations are used hereafter:

10	EDTA	ethylenediaminetetraacetic acid
	IPTG	Isopropyl-β-D-thiogalactopyranoside
	MeCN	methyl cyanide (acetonitrile)
	Mi	Macadamia integrifolia
	MiAMP2	Macadamia integrifolia antimicrobial protein series number 2
15	Ni-NTA	Nickel-nitrilotriacetic acid chromatography media
	ND	not determined
	PCR	polymerase chain reaction
	PMSF	phenylmethylsulphonyl fluoride
	SDS-PAGE	sodium-dodecylsulphate polyacrylamide gel electrophoresis
20	TFA	trifluoroacetate

The term homologue is used herein to denote any polypeptide having substantial similarity in composition and sequence to the polypeptide used as the reference. The homologue of a reference polypeptide will contain key elements such as cysteine or other residues spaced at identical intervals such that a substantially similar three-dimensional global structure is adopted by the homologue as compared to the reference. The homologue will also exhibit substantially the same antimicrobial activity as the reference protein.

The present inventors have identified a new class of proteins with antimicrobial activity. Prototype proteins can be isolated from seeds of *Macadamia integrifolia*. The invention thus provides antimicrobial proteins *per se* and also DNA sequences encoding these antimicrobial proteins.

The invention also provides amino acid sequences of proteins which are homologous to the prototype antimicrobial proteins from *Macadamia integrifolia*. Thus, in addition to the antimicrobial proteins from Macadamia, this invention also provides amino acid sequences of homologues from other species which have hitherto been unrecognized as having antimicrobial activity.

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While the first antimicrobial protein in the present series was isolated directly from Macadamia integrifolia, additional antimicrobial proteins were identified through cloning efforts, homology searches and subsequent antimicrobial testing of the encoded proteins after expression in and purification from liquid culture. After the first protein from this series was purified from macadamia and termed MiAMP2, clones were obtained which encoded a preproprotein containing MiAMP2. This large protein (666 amino acids), represented by several almost identical clones, contained four adjacent regions with significant similarity to the purified antimicrobial protein fragment (MiAMP2) which itself was found to lie within region three in the cloned nucleotide sequence; hence the purified antimicrobial protein is termed MiAMP2c. Other fragments contained in the 666-amino-acid clone are termed MiAMP2a, b and d as per their locations in the cloned nucleotide sequence. Several other sequences with significant homology to the MiAMP2a, b, c, and d protein fragments were then identifed in the Entrez data base. These homologous sequences were contained within larger seed storage proteins from cotton and cocoa which sequences had not been previously described as containing antimicrobial protein sequences or as exhibiting antimicrobial activity. Fragments of larger seed storage proteins containing sequences homologous to MiAMP2c were tested and are here demonstrated to exhibit antimicrobial activity. Thus, the inventors have established a process for obtaining antimicrobial protein fragments from larger seed storage proteins. In the light of these findings, it is evident that fragments of other seed storage proteins containing sequences similar to the proteins described will also exhibit antimicrobial activity.

In particular, the 47-amino-acid TcAMP1 (for *Theobroma cacao* antimicrobial protein 1) and the 60-amino-acid TcAMP2 sequences were derived from a cocoa vicilin seed storage protein gene sequence (which contains 525 amino acids) (Spencer, M.E. and Hodge R. [1992] *Planta* 186:567-576). These derived fragments were then expressed in liquid culture. Cocoa vicilin fragments thus expressed and subsequently purified (Examples 10 and 11), were shown to be antimicrobial (Example 15). This is the first report that fragments of the cocoa vicilin protein possess antimicrobial activity. Pools of sequences containing fragments homologous to the MiAMP2c apparently released from cotton vicilin seed storage protein have been shown to possess antimicrobial activity (Chung, R. P.T. *et al.* [1997] *Plant Science* 127:1-16). This finding is clearly embodied in sequences disclosed in this application.

In addition to showing that cocoa-vicilin-derived fragments exhibit antimicrobial activity, there is herein described additional proteins which exhibit antimicrobial activity. For example, there is described below proteins from *Stenocarpus sinuatus* which are of similar size to MiAMP2 subunits, react with MiAMP2c antiserum, and contain sequences homologous to MiAMP2 proteins (see Figure 4). Based on the evidence provided herein, sequences homologous to the MiAMP2c

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subunit (i.e., MiAMP2a, b, d; TcAMP1; TcAMP2; and cotton fragments 1, 2 and 3—see Figure 4) constitute proteins which contain the fragment with antimicrobial activity. The antimicrobial activity of MiAMP2 fragments from macadamia, and the TcAMP1 and 2 fragments from cocoa, is exemplified below. R. P. T. Chung *et al.* (*Plant Science* 127:1-16 [1997]) have demonstrated that the cotton fragments exhibit antimicrobial activity. Other antimicrobial proteins can also be derived from seed storage proteins such as peanut allergen Ara h (Burks, A.W. *et al.* [1995] *J. Clin. Invest.* 96 (4), 1715-1721), maize globulin (Belanger, F. C. and Kriz, A. L.[1991] *Genetics* 129 (3), 863-872), barley globulin (Heck, G. R. *et al.* [1993] *Mol. Gen. Genet.* 239 (1-2), 209-218), and soybean conglycinin (Sebastiani, F. L. *et al.* [1990] *Plant Mol. Biol.* 15 (1), 197-201), all of which contain the same key elements which are present in the sequences which are here shown to exhibit antimicrobial activity.

The proteins which contain regions of sequence homologous to MiAMP2 (as in Figure 4) can be used to construct nucleotide sequences encoding 1) the active fragments of larger proteins, or 2) fusions of multiple antimicrobial fragments. This can be done using standard codon tables and cloning methods as described in laboratory manuals such as *Current Protocols in Molecular Biology* (copyright 1987-1995 edited by Ausubel F. M. *et al.* and published by John Wiley & Sons, Inc., printed in the USA). Subsequently, these can be expressed in liquid culture for purification and testing, or the sequences can be expressed in transgenic plants after placing them in appropriate expression vectors.

The antimicrobial proteins *per se* will manifest a particular three-dimensional structure which may be determined using X-ray crystallography or nuclear magnetic resonance techniques. This structure will be responsible in large part for the antimicrobial activity of the protein. The sequence of the protein can also be subjected to structure prediction algorithms to assess whether any secondary structure elements are likely to be exhibited by the protein (see Example 8 and Figure 7). Secondary structures, thus predicted, can then be used to model three-dimensional global structures. Although three-dimensional structure prediction is not feasible for most proteins, the secondary structure predictions for MiAMP2c were sufficiently simple and clear that a three-dimensional model structure has been obtained for the MiAMP2c protein. Homologues exhibiting the same cysteine spacing and other key elements will also adopt the same three-dimensional structure. Example 8 shows that the structure most likely to be adopted by MiAMP2c (and homologues) is a helix-turn-helix structure stabilised by at least two disulfide bridges connecting the two antiparallel  $\alpha$ -helical segments (see Figure 8). Additional stabilisation can be provided by an extra disulfide bridge (e.g., as in MiAMP2b) or by a hydrophobic ring-stacking interaction between tyrosine and/or phenylalanine residues (e.g., MiAMP2a and MiAMP2c), each located on the same face of the  $\alpha$ -

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helical segments as the normally present cysteine residues which participate in the 2 disulfide linkages mentioned above. NMR signals exhibited by MiAMP2c are consistent with the three-dimensional global model produced from the secondary-structure predictions mentioned above.

It will be appreciated that one skilled in the art could take a protein with known structure, alter the sequence significantly, and yet retain the overall three-dimensional shape and antimicrobial activity of the protein. One aspect of the structure that most likely could not be altered without seriously affecting the structure (and, therefore, the activity of the protein) is the content and spacing of the cysteine residues since this would disrupt the formation of disulfide bonds which are critical to a) maintaining the overall structure of the protein and/or b) making the protein more resistant to denaturation and proteolysis (stabilizing the protein structure). In particular, it is essential that cysteine residues reside on one face of the helix in which they are contained. This can best be accomplished by maintaining a three-residue spacing between the cysteine residues within each helix, but, can also be accomplished with a two-residue interval between the cysteine residues provided the cysteines on the other helical segment are separated by three residues (i.e., C-X-X-C-X-X-X-C-nX-C-X-X-X-C-X-X-C where C is cysteine, X is any amino acid, and n is the number of residues forming a turn between the two  $\alpha$ -helical segments). Aromatic tyrosine (or phenylalanine) residues can also function to add stability to the protein structure if they are located on the same face of the helix as the cysteine side chains. This can be accomplished by providing appropriate spacing of two or three residues between the aromatic residue and the proximate cysteine residue (i.e., Z-X-X-C-X-X-X-C-nX-C-X-X-X-Z where Z is tyrosine or phenylalanine).

The distribution of positive (and negative) charges on the various surfaces of the protein will also serve a critical role in determining the structure and activity of the protein. In particular, the distribution of positively-charged residues in an  $\alpha$ -helical region of a protein can result in positive charges lying on one face of the helix or may result in the charged residues being concentrated in some particular portion of the molecule. An alternative distribution of positively charged residues is for them to project into the solvent in a radial orientation to the core of the protein. This orientation is predicted for several of the MiAMP2 homologues (data not shown). The spacing which is required for positioning of the residues on one face of the helix or the spacing required to accomplish a radial orientation from the core can easily be determined by one skilled in the art using a helical wheel plot with the sequence of interest. A helical wheel plot uses the fact that, in  $\alpha$ -helices, each turn of the helix is composed of 3.6 residues on average. This number translates to 100° of rotational translation per residue making it possible to construct a plot showing the distribution of side chains in a helical region. Figure 8 shows how the spacing of charged residues can lead to most of the

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positively charged side chains being localised on one face of the helix. It will be appreciated by one of skill in the art that positive charges are conferred by arginine and lysine residues.

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In order for the protein to develop into a helix-turn-helix structure, it is also necessary to have particular residues that favor α-helix formation and that also favor a turn structure in the middle portion of the amino acid sequence (and disfavor a helical structure in the turn region). This can be accomplished by a proline residue or residues in the middle of the turn segment as seen with many of the MiAMP2 homologues. When proline is not present, glycine can also contribute to breaking a continuous helix structure, and inducing the formation of a turn at this position. In one case (i.e., TcAMP1), it appears that serine may be taking on this role. It will be appreciated that the residues in this region of the protein will usually favor the fomation of a turn structure; residues which fulfill this requirement include proline, glycine, serine, and aspartic acid; but, other residues are also allowed.

The DNA sequences reported here are an extremely powerful tool which can be used to obtain homologous genes from other species. Using the DNA sequences, one skilled in the art can design and synthesise oligonucleotide probes which can be used to screen cDNA libraries from other species of plants for the presence of genes encoding antimicrobial proteins homologous to the ones described here. This would simply involve construction of a cDNA library and subsequent screening of the library using as the oligonucleotide probe one or part of one of the sequences reported here (such as sequence ID. No. 2 or the PCR fragment described in Example 9). Other oligonucleotide sequences coding for proteins homologous to MiAMP2 can also be used for this purpose (e.g., DNA sequences corresponding to cotton and cocoa vicilins). Making and screening of a cDNA library can be carried out by purchasing a kit for said purpose (e.g., from Stratagene) or by following well established protocols described in available DNA cloning manuals (see Current Protocols in Molecular Biology, supra). It is relatively straight forward to construct libraries of various species and to specifically isolate vicilin homologues which are similar to the Macadamia, cotton, or cocoa vicilins by using a simple DNA hybridization technique to screen such libraries. Once cloned, these vicilin-related sequences can then be examined for the presence of MiAMP2-like subunits. Such subunits can easily be expressed in E. coli using the system described in Examples 10 and 11. Subsequently, these proteins can also be expressed in transgenic.

Genes, or fragments thereof, under the control of a constitutive or inducible promoter, can then be cloned into a biological system which allows expression of the protein encoded thereby.

Transformation methods allowing for the protein to be expressed in a variety of systems are known. The protein can thus be expressed in any suitable system for the purpose of producing the protein for further use. Suitable hosts for the expression of the protein include *E. coli*, fungal cells, insect cells,

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mammalian cells, and plants. Standard methods for expressing proteins in such hosts are described in a variety of texts including section 16 (Protein Expression) of *Current Protocols in Molecular Biology (supra)*.

Plant cells can be transformed with DNA constructs of the invention according to a variety of known methods (Agrobacterium, Ti plasmids, electroporation, micro-injections, micro-projectile gun, and the like). DNA sequences encoding the Macadamia integrifolia antimicrobial protein subunits (i.e. fragments a, b, c, or d from the MiAMP2 clones) as well as DNA coding for other homologues can be used in conjunction with a DNA sequence encoding a preprotein from which the mature protein is produced. This preprotein can contain a native or synthetic signal peptide sequence which will target the protein to a particular cell compartment (e.g., the apoplast or the vacuole). These coding sequences can be ligated to a plant promoter sequence that will ensure strong expression in plant cells. This promoter sequence might ensure strong constitutive expression of the protein in most or all plant cells, it may be a promoter which ensures expression in specific tissues or cells that are susceptible to microbial infection and it may also be a promoter which ensures strong induction of expression during the infection process. These types of gene cassettes will also include a transcription termination and polyadenylation sequence 3' of the antimicrobial protein coding region to ensure efficient production and stabilisation of the mRNA encoding the antimicrobial proteins. It is possible that efficient expression of the antimicrobial proteins disclosed herein might be facilitated by inclusion of their individual DNA sequences into a sequence encoding a much larger protein which is processed in planta to produce one or more active MiAMP2-like fragments.

Gene cassettes encoding the MiAMP2 series antimicrobial proteins (i.e., MiAMP2a, b, c, or d; or all of the subunits together; or the entire MiAMP2 clone) or homologues of the MiAMP2 proteins as described above can then be expressed in plant cells using two common methods. Firstly, the gene cassettes can be ligated into binary vectors carrying: i) left and right border sequences that flank the T-DNA of the *Agrobacterium tumefaciens* Ti plasmid; ii) a suitable selectable marker gene for the selection of antibiotic resistant plant cells; iii) origins of replication that function in either *A. tumefaciens* or *Escherichia coli*; and iv) antibiotic resistance genes that allow selection of plasmid-carrying cells of *A. tumefaciens* and *E. coli*. This binary vector carrying the chimaeric MiAMP2 encoding gene can be introduced by either electroporation or triparental mating into *A. tumefaciens* strains carrying disarmed Ti plasmids such as strains LBA4404, GV3101, and AGL1 or into *A. rhizogenes* strains such as A4 or NCCP1885. These *Agrobacterium* strains can then be co-cultivated with suitable plant explants or intact plant tissue and the transformed plant cells and/or regenerants selected using antibiotic resistance.

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A second method of gene transfer to plants can be achieved by direct insertion of the gene in target plant cells. For example, an MiAMP2-encoding gene cassette can be co-precipitated onto gold or tungsten particles along with a plasmid encoding a chimaeric gene for antibiotic resistance in plants. The tungsten particles can be accelerated using a fast flow of helium gas and the particles allowed to bombard a suitable plant tissue. This can be an embryogenic cell culture, a plant explant, a callus tissue or cell suspension or an intact meristem. Plants can be recovered using the antibiotic resistance gene for selection and antibodies used to detect plant cells expressing the MiAMP2 proteins or related fragments.

The expression of MiAMP2 proteins in the transgenic plants can be detected using either antibodies raised to the protein(s) or using antimicrobial bioassays. These and other related methods for the expression of MiAMP2 proteins or fragments thereof in plants are described in *Plant Molecular Biology* (2nd ed., edited by Gelvin, S.B. and Schilperoort, R.A., © 1994, published by Kluwer Academic Publishers, Dordrecht, The Netherlands)

Both monocotyledonous and dicotyledonous plants can be transformed and regenerated. Examples of genetically modified plants include maize, banana, peanut, field peas, sunflower, tomato, canola, tobacco, wheat, barley, oats, potato, soybeans, cotton, carnations, roses, sorghum. These, as well as other agricultural plants can be transformed with the antimicrobial genes such that they would exhibit a greater degree of resistance to pathogen attack. Alternatively, the proteins can be used for the control of diseases by topological application.

The invention also relates to application of antimicrobial protein in the control of pathogens of mammals, including humans. The protein can be used either in topological or intravenous applications for the control of microbial infections.

As indicated above in the description of the tenth embodiment, the invention includes within its scope the preparation of antimicrobial proteins based on the prototype MiAMP2 series of proteins. New sequences can be designed from the MiAMP2 amino acid sequences which substantially retain the distribution of positively charged residues relative to cysteine residues as found in the MiAMP2 proteins. The new sequence can be synthesised or expressed from a gene encoding the sequence in an appropriate host cell. Suitable methods for such procedures have been described above. Expression of the new protein in a genetically engineered cell will typically result in a product having a correct three-dimensional structure, including correctly formed disulphide linkages between cysteine residues. However, even if the protein is chemically synthesised, methods are known in the art for further processing of the protein to break undesireable disulfide bridges and form the bridges between the desired cysteine residues to give the desired three-dimensional structure should this be necessary.

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Macadamia integrifolia antimicrobial proteins series number 2

As indicated above, a new series of potent antimicrobial proteins has been identified in the seeds of *Macadamia integrifolia*. The proteins collectivelly are called the MiAMP2 series of antimicrobial proteins (or MiAMP2 proteins) because they are all found on one large preproprotein which is processed into smaller subunits, each exhibiting antimicrobial activity; they represent the second class of antimicrobial proteins isolated from *Macadamia integrifolia*. Each protein fragment of the series has a characteristic pI value. MiAMP2a, b, c, and d subunits as shown in Figure 4 have predicted pI values of 4.4, 4.6, 11.5, and 11.6 respectively (predicted using raw sequence data without the His tag or cleavage sequences associated with expression of fragments in the vector pET16b), and contain two sets of CXXXC motifs which are important in stabilising the three-dimensional structure of the protein through the formation of disulfide bonds. Additionally, the proteins contain either an added set of aromatic (tyrosine/phenylalanine) residues or an added set of cysteine residues located at positions which would give more stability to the helix-turn-helix structure as described above and in Example 8.

The amino acid sequences of the MiAMP2 series of proteins share significant homology with fragments of previously described proteins in sequence databases (Swiss Prot and Non-redundant databases) searched using the BLASTP algorithm (Altschul, S.F. et al. [1990] J. Mol. Biol. 215:403). In particular, MiAMP2a, b, c and d sequences exhibit significant similarity with regions of cocoa vicilin and cotton vicilin (as seen in Figure 6). Some similarity is also seen with fragments from other seed storage proteins of peanut (Burks, A. W. et al. [1995] J. Clin. Invest. 96 (4), 1715-1721), maize (Belanger, F. C. and Kriz, A. L.[1991] Genetics 129 (3), 863-872), barley (Heck, G. R. et al. [1993] Mol. Gen. Genet. 239 (1-2), 209-218), and soybean (Sebastiani, F. L. et al. [1990] Plant Mol. Biol. 15 (1), 197-201). Although, in some cases the homology is not extremely high (for example, 18% identity between MiAMP2a and cotton subunit 1; see Figure 4), the spacing of the main four cysteine residues is conserved in all subunits and homologues. In addition, both cotton and cocoa vicilin-derived subunits retain the conserved tyrosine or phenylalanine residues as additional stabilisers of the tertiary structure. The cotton and cocoa vicilins with 525 and 590 amino acids, respectively, are much larger proteins than MiAMP2c (47 amino acids) (see Figures 4 and 6). Although MiAMP2 subunits also share some homology with MBP-1 antimicrobial protein from maize (Duvick, J.P. et al. (1992) J Biol Chem 267:18814-20) the number of residues between the CXXXC motifs is 13 which puts MBP-1 outside the specifications for the spacing given here in this application. MBP-1 is also a smaller protein (33 amino acids), overall, than the sequences claimed here and there is no evidence available the MBP-1 is derived from a larger seed storage protein other than some similarity with a portion of miaze globulin protein. However, MBP-1 cannot be derived

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from from the maize globulin since maize globulin contains 10 residues between the two CXXXC motifs while MBP-1 contains 13. The alignments in Figures 4 and 6 show the similarity in cysteine spacing between MiAMP2 subunits and the cocoa and cotton vicilin-derived molecules. The cysteine and the aromatic tyrosine/phenylalanine residues in Figures 4 and 6 are highlighted with bold underlined text. Figure 4 also shows the alignment of additional proteins which can be expressed in liquid culture and shown to exhibit antimicrobial activity.

All of the MiAMP2 homologues that have been tested exhibit antifungal activity. MiAMP2 homologues show very significant inhibition of fungal growth at concentrations as low as 2  $\mu$ g/ml for some of the pathogens/microbes against which the proteins were tested. Thus they can be used to provide protection against several plant diseases. MiAMP2 homologues can be used as fungicides or antibiotics by application to plant parts. The proteins can also be used to inhibit growth of pathogens by expressing them in transgenic plants. The proteins can also be used for the control of human pathogens by topological application or intravenous injection. One characteristic of the proteins is that inhibition of some microbes is suppressed by the presence of Ca<sup>2+</sup> (1 mM). An example of this effect is provided for MiAMP2c subunit in Table 1.

Some of the MiAMP2 proteins and homologues could also function as insect control agents. Since some of the proteins are extremely basic (e.g., pI > 11.5 for MiAMP2c and d subunits), they would maintain a strong net-positive charge even in the highly alkaline environment of an insect gut. This strong net-positive charge would enable it to interact with negatively charged structures within the gut. This interaction may lead to inefficient feeding, slowing of growth, and possibly death of the insect pest.

Non-limiting examples of the invention follow.

#### Example 1

## Extraction of Basic Protein from Macadamia integrifolia Seeds

Twenty five kilograms of Mi nuts (purchased from the Macadamia Nut Factory, Queensland, Australia) were ground in a food processor (The Big Oscar, Sunbeam) and the resulting meal was extracted for 2-4 hours at 4°C with 50 L of an ice-cold extraction buffer containing 10 mM NaH2PO4, 15 mM Na2HPO4, 100 mM KCl, 2 mM EDTA, 0.75% polyvinylpolypyrolidone, and 0.5 mM phenylmethylsulfonyl fluoride (PMSF). The resulting homogenate was run through a kitchen strainer to remove larger particulate material and then further clarified by centrifugation (4000 rpm for 15 min) in a large capacity centrifuge. Solid ammonium sulphate was added to the supernatant to obtain 30% relative saturation and the precipitate allowed to form overnight with stirring at 4°C. Following centrifugation at 4000 rpm for 30 min, the supernatant was taken and ammonium sulphate added to achieve 70% relative saturation. The solution was allowed to precipitate overnight and then

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centrifuged at 4000 rpm for 30 min in order to collect the precipitated protein fraction. The precipitated protein was resuspended in a minimal volume of extraction buffer and centrifuged once again (13,000 rpm x 30 min) to remove the any insoluble material yet remaining. After dialysis (10 mM ethanolamine pH 9.0, 2 mM EDTA and 1 mM PMSF) to remove residual ammonium sulphate, the protein solution was passed through a Q-Sepharose Fast Flow column (5 x 12 cm) previously equilibrated with 10 mM ethanolamine (pH 9), 2 mM in EDTA). The collected flowthrough from this column represents the basic (pI >9) protein fraction of the seeds. This fraction was further purified as described in Example 3.

#### Example 2

Antifungal and Antibacterial Activity Assays

In general, bioassays to assess antifungal and antibacterial activity were carried out in 96-well microtitre plates. Typically, the test organism was suspended in a synthetic growth medium consisting of K2HPO4 (2.5 mM), MgSO4 (50 μM), CaCl<sub>2</sub> (50 μM), FeSO4 (5 μM), CoCl<sub>2</sub> (0.1 μM), CuSO4 (0.1 μM), Na<sub>2</sub>MoO4 (2 μM), H<sub>3</sub>BO<sub>3</sub> (0.5 μM), KI (0.1 μM), ZnSO4 (0.5 μM), MnSO4 (0.1 μM), glucose (10 g/L), asparagine (1 g/L), methionine (20 mg/L), myo-inositol (2 mg/L), biotin (0.2 mg/L), thiamine-HCl (1 mg/L) and pyridoxine-HCL (0.2 mg/L). The test organism consisted of bacterial cells, fungal spores (50,000 spores/ml) or fungal mycelial fragments (produced by blending a hyphal mass from a culture of the fungus to be tested and then filtering through a fine mesh to remove larger hyphal masses). Fifty microlitres of the test organism suspended in medium was placed into each well of the microtitre plate. A further 50 μl of the test antimicrobial solution was added to appropriate wells. To deal with well-to-well variability in the bioassay, 4 replicates of each test solution were done. Sixteen wells from each 96-well plate were used as controls for comparison with the test solutions.

Unless otherwise stated, incubation was at 25°C for 48 hours. All fungi including yeast were grown at 25°C. *E. coli* were grown at 37°C and other bacteria were bioassayed at 28°C. Percent growth inhibition was measured by following the absorbance at 600 nm of growing cultures over various time intervals and is defined as 100 times the ratio of the average change in absorbance in the control wells minus the change in absorbance in the test well divided by the average change in absorbance at 600 nm for the control wells (i.e., [(avg change in control wells - change in test well) / (avg change in control wells)] x 100). Typically, measurements were taken at 24 hour intervals and the period from 24-48 hours was used for %Inhibition measurements.

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#### Example 3

Purification of antimicrobial protein from Macadamia integrifolia basic protein fraction The starting material for the isolation of the Mi antimicrobial protein was the basic fraction extracted from the mature seeds as described above in Example 1. This protein was further purified by cation exchange chromatography as shown in Figure 1.

About 4 g of the basic protein fraction dissolved in 20 mM sodium succinate (pH 4) was applied to an S-Sepharose High Performance column (5 X 60 cm) (Pharmacia) previously equilibrated with the succinate buffer. The column was eluted at 17 ml/min with a linear gradient of 20 L from 0 to 2 M NaCl in 20 mM sodium succinate (pH 4). The cluate was monitored for protein by on-line measurement of the absorbance at 280 nm and collected in 200 ml fractions. Portions of each fraction were subsequently tested in the antifungal activity assay against Phytopthora cryptogea at a concentration of 100 µg/ml in the presence and absence of 1 mM Ca<sup>2+</sup>. Results of bioassays are included in Figures 1a and 1b where the elution gradient is shown as a solid line and the shaded bars represent %Inhibition. The Figure 1a assays were conducted without added Ca<sup>2+</sup> while 1 mM Ca<sup>2+</sup> was included in the Figure 1b assays. Fractionation yielded a number of unresolved peaks eluting between 0.05 and 2 M NaCl. A peak eluting at about 16 hours into the separation (fractions 90-92) showed significant antimicrobial activity.

Fractions showing significant antimicrobial activity were further purified by reversed-phase chromatography. Aliquots of fractions 90-92 were loaded onto a Pep-S (C2/C18), column (25 x 0.93 cm) (Pharmacia) equilibrated with 95% H<sub>2</sub>O/5% MeCN/0.1% TFA (=100%A). The column was eluted at 3 ml/min with a 240 ml linear gradient (80 min) from 100%A to 100%B (=5% H<sub>2</sub>O/95% MeCN/0.1% TFA). Individual peaks were collected, vacuum dried three times in order to remove traces of TFA, and subsequently resuspended in 500 microlitres of milli-Q water (Millipore Corporation water purification system) for use in bioassays as described in Example 2. Figure 2 shows the HPLC profile of purified fraction 92 from the cation-exchange separation shown in Figures 1 and 2. Protein elution was monitored at 214 nm. The acetonitrile gradient is shown by the straight line. Individual peaks were bioassayed for antimicrobial activity: the bars in Figure 3 show the inhibition corresponding to 15 μg/ml of material from each of the fractions. The active protein elutes at approximately 27 min (~30% MeCN/0.1%TFA) and is called MiAMP2c.

Example 4

#### Purity of Isolated MiAMP2c

The purity of the isolated antimicrobial protein was verified by native SDS-PAGE followed by staining with coomassie blue protein staining solution. Electrophoresis was performed on a 10-20% tricine gradient gel (Novex) as per the manufacturers recommendations (100 V, 1-2 hour separation

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time). Under these conditions the purified MiAMP2c migrates as a single discrete band (<10 kDa in size). The detection of a single major band in the SDS-PAGE analysis together with single peaks eluting in the cation-exchange and reversed-phase separations (not shown), gives strong indication that the MiAMP2c preparation is greater than 95% pure and therefore the activity of the preparation was almost certainly due to the MiAMP2c alone and not to a minor contaminating component. A clean signal in mass spectrometric analysis (Example 5 below) also supports this conclusion.

#### Example 5

#### Mass Spectroscopic Analysis of MiAMP2c

Purified MiAMP2c was submitted for mass spectroscopic analysis. Approximately 1 µg of protein in solution was used for testing. Analysis showed the protein to have a molecular weight of 6216.8 Da ± 2 Da. Additionally, the protein was subjected to reduction of disulfide bonds with dithiothreitol and alkylation with 4-vinylpyridine. The product of this reduction/alkylation was then submitted for mass spectroscopic analysis and was shown to have gained 427 mass units (i.e. molecular weight was increased by approximately 4 X 106 Da). The gain in mass indicated that four 4-vinylpyridine groups had reacted with the reduced protein, demonstrating that the protein contains a total of 4 cysteine residues. The cysteine content has also been subsequently confirmed through amino acid sequencing.

#### Example 6

## Amino Acid Sequence of MiAMP2c Protein

Approximately 1 µg of the pure protein which had been reduced and alkylated was subjected to Automated Edman degradation N-terminal sequencing. In the first sequencing run, the sequence of the first 39 residues was determined. Subsequently, approximately 1 mg of MiAMP2c was reacted with Cyanogen Bromide which cleaved the protein on the C-terminal side of Methionine-26. The C-terminal fragment generated by the cleavage reaction was purified by reversed-phase HPLC and sequenced, yielding the remaining sequence of MiAMP2c (i.e. residues 27-47). The full amino acid sequence is RQRDP QQQYE QCQER CQRHE TEPRH MQTCQ QRCER RYEKE KRKQQ KR and represents amino acids 118 to 164 of clone 3 from Example 9 (see Figure 6 and SEQUENCE ID NO: 5). In the figure, cysteine residues are in bold type and underlined to facilitate recognition of the spacing patterns. Depending on the number of disulfide bonds that are formed, the protein mass will range from 6215.6 to 6219.6 Da. This is in close agreement with the mass of 6216.8 ± 2 Da obtained by mass spectrometric analysis (Example 5). The measured mass closely approximates the predicted mass of MiAMP2c in a two-disulfide form as is expected to be the case.

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#### Example 7

Synthetic DNA Sequence Coding for MiAMP2c with a leader peptide

Using standard codon tables it is possible to reverse-translate the protein sequences to obtain DNA sequences that will code for the antimicrobial proteins. The software program MacVector 4.5.3 was used to enter the protein sequence and obtain a degenerate nucleotide sequence. A codon usage table for tobacco was referenced in order to pick codons that would be adequately represented in tobacco for purposes of obtaining high expression in this test plant. A 30 amino-acid leader peptide was also designed to ensure efficient processing of the signal peptide and secretion of the peptide extracellularly. For this purpose, the method of Von Hiejne was used to evaluate a series of possible leader sequences for probability of cleavage at the correct position [Von Hiejne, G.(1986) Nucleic Acids Research 14(11): 4683-4690]. In particular, the amino acid sequence MAWFH VSVCN AVFVV IIIIM LLMFV PVVRG (Sequence ID. No. 11) was found to give an optimal probability of correct processing of the signal peptide immediately following the G (Gly) of this leader sequence. A 5' untranslated region from tobacco mosaic virus was also added to this synthetic gene to promote higher translational efficiency [Dowson, M.J., et al. (1994) Plant Mol. Biol. Rep. 12(4):347-357]. The synthetic gene also contains restriction sites at the 5' and 3' ends and immediately 5' of the start ATG for efficient cloning and subcloning procedures. Figure 5 shows a synthetic DNA sequence suitable for use in plant expression experiments. In this Figure, the arrow shows where translation is initiated and the triangular symbol indicates the point of cleavage of the signal peptide.

#### Example 8

#### Structure prediction of MiAMP2c Protein

Using sequence analysis algorithms, putative secondary structure motifs can be assigned to the protein. Five different algorithms were used to predict whether α-helices, β-sheets, or turns can occur in the MiAMP2c protein (Figure 4). Methods were obtained from the following sources: DPM method, Deleage, G., and Roux, B. (1987) *Prot. Eng.* 1:289-294; SOPMA method, Geourjon, C., and Deleage, G. (1994) *Prot. Eng.* 7:157-164; Gibrat method, Gibrat, J.F., Garnier, J., and Robson, B.(1987) *J.Mol.Biol.* 198:425-443; Levin method, Levin, J.M., Robson, B., and Garnier, J. (1986) *FEBS Lett.* 205:303-308; and PhD method, Rost, B., And Sander, C. (1994) *Proteins* 19:55-72. Figure 7 shows the predicted locations of α-helices, β-sheets and turns. The following symbols have been used in Figure 7: C, coil (unstructured); H, alpha helix; E, β- sheet; and S, turn. Underlined residues are those which were predicted to exhibit an α-helical structure by at least 2 separate structure prediction methods; these are represented as helices in Figure 8.

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It is clear from the secondary structure predictions that the protein is highly  $\alpha$ -helical. While secondary structure prediction is often difficult and inaccurate, this particular prediction gives a clear indication of the structure of the protein. Examination of the secondary-structure predictions show a clear preponderance of two  $\alpha$ -helical regions broken by a stretch of about 5-8 residues. This is highly suggestive of a helix-turn-helix motif.

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Helical wheel analysis of the MiAMP2c amino acid sequence shows that cysteine residues with a CXXXC spacing will be aligned on one face of the helix in which they are located. Since the cysteines are involved in disulfide bond formation, the cysteine side chains in one helix must form covalent bonds with the cysteine side chains located on the other helical segment. When the helical segments are arranged in such a way as to bring the cysteine side chains from each respective helix into proximity with the other cysteine side chains, the resulting three-dimensional structure is shown in Figure 8. This structure exhibits a remarkable distribution of positively charge residues on one face of the protein comprised of two helices held together by two disulfide bonds. Figure 8 shows how the spacing of positively charged residues in helical regions of this molecule will cause these side chains to lie on one face of the helix. The positively charged residues are the dark side chains outlined in black. Other dark side chains represent acidic residues. A proline residue (grey colour marked with a 'P') is located at the extreme left end of the molecule in the turn region. Solid black lines show where disulfide bonds connect the two helices. The dotted line shows where the two aromatic hydrophobic residues interact to add stability to the helix-turn-helix structure.

This helix-turn-helix structure will be adopted by all MiAMP2 homologues containing the same cysteine spacing and residues with helix and turn-forming propensities. Other MiAMP2 fragment sequences can be superimposed onto the global structure shown in figure 8. The overall structure will remain essentially the same but the charge distribution will vary according to the sequences involved. In the case of MiAMP2b, the dotted line would represent an added disulfide bridge instead of a hydrophobic interaction.

#### Example 9

cDNA cloning of genes corresponding to MiAMP2c

## PCR Amplification of a genomic fragment of the MiAMP2c gene

Using the reverse-translated nucleotide sequences, degenerate primers were made for use in PCR reactions with genomic DNA from Macadamia. Primer JPM17 sequence was 5' CAG CAG CAG CAG TAT GAG CAG TG 3' and primer JPM20 degenerate sequence was 5' TTT TTC GTA (T/T)C(T/G) (G/T)C(T/G) TTC GCA 3' (SEQ ID NOS: 12 and 13). Primers JPM17 and JPM20 were used in PCR amplifications carried out for 30 cycles with 30 sec at 95°C, 1 min at 50°C, and 1 min at 72°C. PCR products with sizes close to those which were expected were directly sequenced

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(ABI PRISM Dye Terminator Cycle Sequencing Ready Reaction Kit from Perkin Elmer Corporation) after excising DNA bands from agarose gels and purifying them using a Qiagen DNA clean-up kit. Using this approach, it was possible to amplify a fragment of DNA of approximately 100 bp. Direct sequencing of this nucleotide fragment yielded the nucleotide sequence corresponding to a portion of the amino acid sequence of the antimicrobial protein MiAMP2c (amino acids 7-39 of Figure 4). The partial nucleotide sequence obtained from the above-mentioned fragment excluding the primer sequences was 5' TCA GAA GCG CTG CCA ACG GCG CGA GAC AGA GCC ACG ACA CAT GCA AAT TTG TCA ACA ACG C 3' (corresponding to base pairs 264 to 324 in SEQ ID NO: 6). This sequence can be used for a variety of purposes including screening of cDNA and genomic libraries for clones of MiAMP2 homologues or design of specific primers for PCR amplification reactions.

#### Messenger RNA isolation from Macadamia nut kernels

Fifty-eight grams of Macadamia nut kernels were ground to powder under liquid nitrogen using a mortar and pestel. RNA from ground material was then purified using a Guanidine thiocyanate/Cesium chloride technique (*Current Protocols in Molecular Biology, supra*). Using this method approximately 5 mg of total RNA was isolated. Messenger RNA was then purified from total RNA using a spun column mRNA purification kit (Pharmacia).

#### cDNA library construction

A cDNA library was constructed in a lambda ZAP vector using a library kit from Stratagene. A total of 6 reactions were performed using 25 micrograms of messenger RNA. First and second strand cDNA synthesis was performed using MMLV Reverse transcriptase and DNA Polymerase I, respectively. After blunting the cDNA with *Pfu* DNA Polymerase, *Eco* RI linker adapters were ligated to the DNA. DNA was then kinased using T4 polynucleotide kinase and the DNA subsequently digested with *Xho* I restriction endonuclease. At this point cDNA material was fractionated according to size using a sephacryl-S500 column supplied with the kit. DNA was then ligated into the lambda ZAP vector. The vector containing ligated insert was then packaged into lambda phage (Gigapack III packaging extract from Stratagene).

#### Screening of library

The library constructed above was then plated and screened in XL1-blue *E.coli* bacterial lawns growing in top agarose. Plaques containing individual clones were isolated by lifting onto Hybond N+ membranes (Amersham LIFE SCIENCE), hybridizing to a radiolabeled version of the genomic DNA fragment amplified above, imaging of the blot, and picking of possitive clones for the next round of screeing. After secondary and tertiary screening, plaques were sufficiently isolated to allow

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picking of single clones. Several clones were obtained, and subsequently the pBK-CMV vector portion from the larger lambda vector was excised.

#### Sequence of MiAMP2c cDNA clones

Vectors (pBK-CMV) containing putative MiAMP2c clones were sequenced to obtain the DNA sequence of the cloned inserts. Seven clones were partially sequenced and an additional three clones were fully sequenced (see SEQ ID NOS: 2, 4 and 6 for DNA sequences of the macadamia clones). Translation of the DNA sequences showed that the full length clones encoded highly similar proteins of 666 amino acids. Figure 6 shows that these proteins have substantial similarity to vicilin seed-strorage proteins from cocoa and cotton. Stars show positions of conserved identities and dots show positions of conserved similarities. Examination of the protein sequences revealed that the exact MiAMP2c sequence is found within the translated protein sequence of clone 3 at amino acid positions 118 to 164 (see Figure 6); clones 1 and 2 contained sequences differing from MiAMP2c by 2 residues and 3 residues, respectively, out of 47 amino acids total in the MiAMP2c sequence.

The translation products of the full-length clones (i.e., clones 1 and 2) consist of a short signal peptide from residues 1 to 28, a hydrophilic region from residues 29 to ~246, and then two segments stretching from residues ~246 to 666 with a stretch of acidic residues separating them at positions 542-546.

Significantly, the hydrophilic region containing the sequence for MiAMP2c, also contains 3 additional segments which are very similar to MiAMP2 (termed MiAMP2a, b and d). These 4 segments (found between residues 28 and ~246) are separated by stretches in which approximately four out of five residues are acidic (usually glutamic acid). These acidic stretches occur at positions 64-68, 111-115, 171-174, and 241-246 and appear to delineate processing sites for cleavage of the 666-residue preproprotein into smaller functional fragments (acidic stretches delineating cleavage sites are shown as bold characters in Figure 6). All four MiAMP2-like segments of the protein contain 2 doublets of cysteine residues separated by 10-12 residues to give the following pattern C-X-X-X-C-(10-12X)-C-X-X-X-C where X is any amino acid, and C is cysteine. All four segments are expected to form helix-turn-helix motifs as decribed in Example 8 above. It is clear that the cysteines in these locations will form disulfide bridges that stabilize the structure of the proteins by holding the two helical portions together.

The predicted helix-turn-helix motifs can be further stabilized in several ways. The first method of stabilization is exemplified in segments 1 and 3 (i.e., residues 29-63 and 118-170, respectively, of the 666-residue Macadamia vicilin-like protein). These segments is the are stabilized by a hydrophobic ring-stacking interaction between two aromatic residues (one on each  $\alpha$ -helical segment); this is normally accomplished with tyrosine residues but phenylalanine is also

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used. As with the cysteine residues, the location of these aromatic residues in the predicted  $\alpha$ -helical segments is critical if they are to offer stabilization to the helix-turn-helix structure. In segments 1 and 3, the aromatic residues are 2 and 3 residues removed from the cysteine doublets as shown here: Z-X-X-C-X-X-C-(10-12X)-C-X-X-X-C-X-X-Z-Where C is cysteine and Z is usually tyrosine but can be substituted with phenylalanine as is done in segment 1.

The second way to stabilize the helix-turn-helix fragment is by using an added disulfide bridge as seen in fragment 2 (residues 71-110). This is accomplished by placing additional cysteine residues 2 and 3 residues removed from the cysteine doublets as shown here: nX-C-X-X-C-X-X-X-C-(10-12X)-C-X-X-X-C-X-X-X-C-nX. This is the only report that the inventors know of where a helix-turn-helix domain in an antimicrobial protein is stabilized by three disulfide bridges. While segment 4 (residues 175-241) does not contain the extra disulfide bridge or the hydrophobic ring-stacking stabilization, it is probably stabilized by means of weaker ionic and or hydrogen bonding interactions.

## Example 10

Vectors for liquid culture expression of MiAMP2 and homologues

PCR primers flanking the nucleotide region coding for MiAMP2c were engineered to contain restriction sites for *Nde* I and *Bam* HI (corresponding to the 5' and 3' ends of the coding region, respectively; Primer JPM31 sequence: 5' A CAC CAT ATG CGA CAA CGT GAT CC 3'; Primer JPM32 sequence: 3' C GTT GTT TTC TCT ATT CCT AGG GTT G 5', SEQ ID NOS: 14 and 15). These primers were then used to amplify the coding region of MiAMP2c DNA. The PCR product from this amplification was then digested with *Nde* I and *Bam* HI and ligated into a pET17b vector (Novagen / Studier, F. W. *et al.* [1986] *J. Mol. Biol.* 189:113) with the coding region in-frame to produce the vector pET17-MiAMP2c.

A similar approach to the one above was used to construct vectors carrying the coding sequences of MiAMP2c homologues (i.e. MiAMP2a, b, and d as well as Tc AMP1, and TcAMP2). To construct the expression vectors for fragments a, b and d in MiAMP2 clone 1, specific PCR primers incorporating the *Nde* I and *Bam* HI sites were designed to amplify the fragments of interest. The products were then digested with the appropriate restriction enzymes and ligated into the *Nde* I/Bam HI sites of a pET16b vector [Novagen] containing a His tag and a Factor Xa cleavage site (amino acid sequence MGHHH HHHHH HHSSG HIEGR HM, SEQ ID NO: 16). The protein products expressed from the pET16b vector is a fusion to the antimicrobial protein. The coding sequences for MiAMP2-like subunits from cocoa (Figure 4, TcAMP1 and TcAMP2) were obtained from the published DNA sequence of the cocoa vicilin gene (Spencer, M. E. and Hodge R. [1992] *Planta* 186:567-576). Two MiAMP2-like fragments within the cocoa vicilin gene were located at

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the 5' end (corresponding to the residues shown in Figure 4), and two sets of complimentary oligonucleotides corresponding to the desired coding sequences were designed. The complimentary oligonucleotides (90 to  $\sim$ 100 bases) corresponding to each cocoa subunit contained a 20bp overlap and also contained the *Nde* I and *Bam* HI restriction endonuclease cut sites.

ì	5	For TcAMP, the following nucleotides were synthesised:				
ons and that the thin and the fan fan flam that the		TcAMP1 forward oligo	5' GGGAATTCCA TATGTATGAG CGTGATCCTC			
			GACAGCAATA CGAGCAATGC CAGAGGCGAT			
			GCGAGTCGGA AGCGACTGAA GAAAGGGAGC 3';			
		TcAMP1 reverse oligo	5' GAAGCGACTG AAGAAAGGGA GCAAGAGCAG			
	10		TGTGAACAAC GCTGTGAAAG GGAGTACAAG			
			GAGCAGCAGA GACAGCAATA GGGATCCACA C 3'.			
		For TcAMP2, the following oligonucleotides were used:				
		TcAMP2 forward oligo	5' GGGAATTCCA TATGCTTCAA AGGCAATACC			
			AGCAATGTCA AGGGCGTTGT CAAGAGCAAC			
	15		AACAGGGCA GAGAGAGCAG CAGCAGTGCC			
			AGAGAAAATG C 3';			
		TcAMP2 reverse oligo	5' GTGTGGATCC CTAGCTCCTA TTTTTTTGT			
			GATTATGGTA ATTCTCGTGC TCGCCTCTCT			
			CTTGTTCCTT ATATTGCTCC CAGCATTTTC			
9; 8;	20		TCTGGCACTG CT 3'.			

The oligonucleotide sets were added to individual PCR amplification reactions in order make individual PCR fragments containing the desired coding region. Since initial PCR amplifications gave fuzzy bands, reamplification of the original products was carried out using new 20mer primers (complimentary to the 5'ends of the forward and reverse oligonucleotides shown above) designed to amplify the entire coding region of the cocoa subunits. Once amplified, the PCR products were restriction digested with the appropriate enzymes and ligated into the vector pET16b as above. This procedure was carried out for both cocoa fragments with similarities to MiAMP2c (shown in Figure 4).

## Example 11

Expression in *E.coli* and purification of MiAMP2c and homologues

Starter cultures (50 ml) of *E.coli* strain BL21 (Grodberg, J. [1988] *J. Bacteriol*. 170:1245) transformed with the appropriate pET construct (Example 10) were added to 500ml of NZCYM media (*Current Protocols in Molecular Biology, supra*) and cultured to an optical density of 0.6 (600 nm) and induced with the addition of 0.4 or 1.0 mM IPTG depending on whether pET17b

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(containing a T7 promoter) or pET16b (containing a His tag fusion and a T7 promoter/lac operator) vector was being used. After cells were induced, cultures were allowed to grow for 4 hours before harvesting. Aliquots of the growing cultures were removed at timed intervals and protein extracts run on an SDS-PAGE gel to follow the expression levels of MiAMP2 and homologues in the cultures. Fragments being expressed with a Histidine tag (i.e., in the pET16b vector), were harvested by centrifuging induced cell cultures at 5000g for 10 minutes. Cell pellets were resuspended and broken by stirring for one hour in 6 M Guanidine-HCl, buffered with 100 mM sodium phosphate and 10 mM Tris at pH 8.0. Broken cell suspensions were centrifuged at 10,000g for 20-30 minutes to settle the cellular debris. Supernatants were removed to fresh tubes and 500 mg of Ni-NTA fast flow resin (Qiagen) was added to each supternatant. After gentle mixing at 4°C for 30-60 minutes, the suspension was loaded into a small column, rinsed two times with 8 M Urea (pH 8.0 and then pH 6.3) and subsequently, the protein was eluted using 8 M Urea pH 4.5. Protein fractions thus obtained were substantially pure but were further purified using an 9.3 x 250 mm C2/C18 reverse phase column (Pharmacia) and 75 minute gradient from 5% to 50% acetonitrile (0.1% TFA) flowing at 3 ml/min (data not shown).

All of the MiAMP2c homologues (except MiAMP2c which was expressed in pET17b) were expressed in the pET16b vector containing the Histidine tag. While induction of the MiAMP2c culture proceded as above, the rest of the purification was somewhat different. In this case, MiAMP2c-expressing cells were harvested by centrifugation but were then resuspended in phosphate buffer (100 mM, pH 7.0 containing 10 mM EDTA and 1 mM PMSF) and broken open using a French press instrument. Cellular debris containing MiAMP2c inclusion bodies was solubilized using a 6 M Guanidine-HCl, 10 mM MES pH 6.0 buffer. Soluble material was then recovered after centrifugation to remove insoluble debris remaining from the solubilization step. Guanidine-HCl soluble material was then dialyzed against 10 mM MES pH 6.0 containing PMSF (1 mM) and EDTA (10 mM). Cation-exchange fractionation was carried out as described in Example 3 except on a smaller scale after the dialysis step. Subsequently, the major eluting protein from the cation-exchange column, which was MiAMP2c, was then further purified using reverse phase HPLC as described in Example 3.

Figure 9 shows the SDS-PAGE gel analysis of the various purification stages obtained following induction with IPTG and subsequent purification of expressed proteins. Samples analysed during the TcAMP1 purification were are as follows: lane 1, molecular weight markers; lane 2, Ni-NTA non-binding fraction; lane 3, rinse of Ni-NTA resin with pH 8 urea; lane 4, rinse of Ni-NTA resin with pH 6.3 urea; lane 5, elution of TcAMP1 with pH 4.5 urea; and lane 6, second elution of TcAMP1 with pH 4.5 urea. TcAMP2 was purified in a similar manner and was also subjected to

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reverse-phase HPLC to further purify the fraction eluting from the Ni-NTA resin. Figure 10 shows the reverse phase purification of cocoa subunit number 2 (TcAMP2).

SDS-PAGE gel analysis of the MiAMP2a, b, and d fragment purification is shown in the second panel of Figure 9. Lane contents are as follows: lane 1, molecular weight markers; lane 2, MiAMP2a pre-induced cellular extractp; lane 3, MiAMP2a IPTG induced cellular extract; lane 4, MiAMP2a Ni-NTA non-binding fraction; lane 5, MiAMP2a elution from Ni-NTA; lane 6, MiAMP2b pre-induced cellular extract; lane 7, MiAMP2b IPTG induced cellular extract; lane 8, MiAMP2b Ni-NTA non-binding fraction; lane 9, MiAMP2b elution from Ni-NTA; lane 10, MiAMP2d pre-induced cellular extract; lane 11, MiAMP2d IPTG induced cellular extract; lane 12, MiAMP2d Ni-NTA non-binding fraction; and lane 13, MiAMP2d elution from Ni-NTA.

Using the vectors described in Example 10, MiAMP2c, and 5 homologues (i.e., MiAMP2a, MiAMP2b, MiAMP2d, TcAMP1 and TcAMP2) were all expressed, purified and tested for antimicrobial activity. The approach taken above can be applied to all of the antimicrobial fragments described in Figure 4. Purified fragments can then be tested for specific inhibition agains microbial pathogens of interest.

#### Example 12

Detection of MiAMP2 homologues in other species using antibodies raised to MiAMP2c Rabbits were immunised intramuscularly according to standard protocols with MiAMP2 conjugated to diphtheria toxoid suspended in Fruends incomplete adjuvent. Serum was harvested from the animals at regular intervals after giving the animal added doses of MiAMP2 adjuvent to boost the immune response. Approximately 100 ml of serum were collected and used for screening of crude extracts obtained from several plant seeds. One hundred gram quantities of seeds were ground and extracted to obtain a crude extract as in Example 1. Aliquots of protein were separated on SDS-PAGE gels and the gels were then blotted onto nitrocellulose membrane for subsequent detection of antibody reacting proteins. The membranes were incubated with MiAMP2c rabbit primary antibodies, washed and then incubated with alkaline phosphatase-conjugated goat anti-rabbit IgG for colorimetric detection of antigenic bands using the chemical 5-bromo-4-chloro-3-indolyl phosphate / nitroblue tetrazolium substrate system (Schleicher and Schuell). Figure 11 shows that various other species contain immunologically-related proteins of similar size to MiAMP2c. Lanes 1-15 contain the extracts from the following species: 1) Stenocarpus sinuatus, 2) Stenocarpus sinuatus(1/10 loading), 3) Restio tremulus, 4) Mesomalaena tetragona, 5) Nitraria billardieri, 6) Petrophile canescens, 7) Synaphae acutiloba, 8) Dryandra formosa, 9) Lambertia inermis, 10) Stirlingia latifolia, 11) Xylomelum angustifolium, 12) Conospermum bracteosum, 13) Conospermum triplinernium, 14) Molecular weight marker, 15) Macacamia integrifolia pure MiAMP2c. Lanes 1-

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13 contain a variety of species, some of which show the presence of antigenically related proteins of a similar size to MiAMP2c. Other bands exhibiting higher molecular weights probably represent the larger precursor seed storage proteins from which the antimcrobial proteins are derived.

Antigenically-related proteins can be seen in lanes 1, 2, 4, 6, 7, 8, 9, and 11-13.

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Bioassays were also performed using crude extracts from various Proteaceae species. Specifically, extracts from *Banksia robur*, *Banksia canei*, *Hakea gibbosa*, *Stenocarpus sinuatus*, and *Stirlingia latifolia* have all been shown to exhibit antimicrobial activity. This activity may derive from MiAMP2 homologues since these species are related to Macadamia.

#### Example 13

Purification of MiAMP2c homologues in another species using antibodies raised to MiAMP2c Based on the detection of immunologically related proteins in other species of the family Proteaceae and the presence of antimicrobial activity in crude extracts, *Stenocarpus sinuatis* was chosen for a large scale fractionation experiment in an attempt to isolate MiAMP2c homologues. Five kg of *S.sinuatus* seed was frozen in liquid nitrogen and ground in a food processor (Big Oscaar Sunbeam). The ground seed was immediately placed into 12 L of 50 mM H<sub>2</sub>SO<sub>4</sub> extraction buffer and extracted at 4°C for 1 hour with stirring. The slurry was then centrifuged for 20 min at 10,000 g to remove particulate matter. The supernatant was then adjusted to pH 9 using a 50mM ammonia solution. PMSF and EDTA were added to final concentrations of 1 and 10 mM respectively.

The crude protein extract was applied to an anion exchange column (Amberlite IRA-938, Rohm and Haas) (3cmx90cm) equilibrated with 50 mM NH4Ac pH 9.0 at a flow rate of 40 ml/min. The unbound protein comprising the basic protein fraction was collected and used in the subsequent purification steps.

The basic protein fraction was adjusted to pH 5.5 with acetic acid and then applied at 10 ml/minute over 12 h to a SP-Sepharose Fast Flow (Pharmacia) Column (5cm x 60cm) preequilibrated with 25mM ammonium acetate. The column was then washed for 3.5 h with 25 mM Acetate pH 5.5. Elution of bound proteins was achieved by applying a linear gradient of NH4Ac from 25 mM to 2.0 M (pH 5.5) at 10 ml/min over 10 h. Absorbance of the eluate was observed at 280 nm and 100 ml fractions collected (see Figure 12).

Cation-exchange fractions that cross-reacted with the antiserum (fractions 14-28, Figure 12) were then further purified by reverse phase chromatography. Cross-reacting fractions were loaded onto a 7 µm C18 reverse phase column (Brownlee) equilibrated with 90% H<sub>2</sub>0, 10% acetonitrile and 0.1% Trifluoroacetic acid (TFA)(=100%A). Bound proteins were eluted with a linear gradient from 100%A to 100%B (5% H<sub>2</sub>0, 95% acetonitrile, 0.08% TFA). The absorbance of the eluted proteins was monitored at 214nm and 280nm. The eluted proteins were dried under vacuum and resuspended

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in water three times to remove traces of TFA from the samples. Reverse phase protein elution fractions 20 to 61 were analysed by pooling 2 adjacent fractions and performing a western blot analysis (see Figure 13). Fractions 22-41 gave a weak positive reaction and fractions 42-57 gave a strong positive reaction to the anti-MiAMP2c antiserum. Fractions that showed antifungal activity against *S.sclerotiorum* at 50  $\mu$ g/ml and 10  $\mu$ g/ml are indicated by arrows on the chromatogram.

Using the approach above, several active fractions (termed SsAMP1 and SsAMP2) were obtained which were assessed for their antifungal activity against *Sclerotinia sclerotiorum*, *Alternaria brassicola*, *Leptosphaeria maculans*, *Verticilium dahliae* and *Fusarium oxysporum*. Bioassays were carried out as described in Example 2 and results shown in Example 15. Another fragment which reacted with MiAMP2 antiserum was purified and sequenced (SsAMP3) but insufficient protein was available for characterisation of antimicrobial activity. Partial sequences obtained from these proteins are shown in Figure 4 (SEQ ID NOS: 26, 27 and 28). Full sequencing of the peptides or cloning of cDNAs encoding the seed storage proteins from this species will reveal the extent of homology between these peptides and MiAMP2-series homologues.

#### Example 14

#### Synthesis of small fragments of MiAMP2c

In an effort to determine if the full MiAMP2c molecule was absolutely necessary for the protein to exhibit antimicrobial activity, two separate peptides were chemically synthesized by Auspep Pty. Ltd. (Australia). For each peptide, the cysteine residues were changed to alanine residues so that disulfide bonds were no longer capable of being formed between two separate protein chains. Tyrosine residues were also changed to alanine since it was expected that tyrosine also participated in the helix-turn-helix stabilization and this would not be needed in the synthetic peptides lacking one of the helices. Alanine is also favorable to the formation of alpha-helices so it should not interfere with the native helical structure to a large degree. Peptide one is comprised of 22 amino acids from 118 to 139 in the amino acid sequence of clone 3 (sequence: RQRDP QQQAE QAQKR AQRRE TE, SEQUENCE ID NO: 9). Peptide 2 is 25 amino acids in length and runs from 140 to 164 in clone 3 (sequence: PRHMQ IAQQR AERRA EKEKR KQQKR, SEQ ID NO: 10). Peptides 1 and 2 are labeled MiAMP2c pep1 and MiAMP2c pep2 respectively. These peptides were resuspended in Milli-Q water and bioassayed against a number of fungi. As seen in Table 2, peptide 2 has inhibitory activity against a variety of fungi whereas peptide 1 exhibited little or no activity. Mixtures of peptide 1 and peptide 2 exhibit similar levels of activity as seen with peptide 2 alone indicating that only peptide 2 is exhibiting activity. The fact that peptide 2 exhibits antimicrobial activity in the absence of the helix-turn-helix structure exhibited by MiAMP2c reveals that the helixturn-helix structure is not absolutely necessary for the peptides to retain activity. Nevertheless,

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peptide 2 did not exhibit the same degree of activity on a molar basis as MiAMP2c (whole fragment) indicating that the helix-turn-helix structure is important for maximal expression of antimicrobial activity by the fragments involved. It is also expected that the helix-turn-helix structure will confer greater stability to the MiAMP2 homologues, thus rendering these proteins less susceptible to proteolytic cleavage and other forms of degredation. Greater stability would lead to maintaining antimicrobial activity over a longer period of time.

# Example 15 Antifungal activity of MiAMP2c homologues and fragment(s)

MiAMP2c and each of the various MiAMP2 homologues were tested against a variety of fungi as concentrations ranging from 2 to 50  $\mu$ g/ml. Table 1 shows the IC50 value of pure MiAMP2c against various fungi and bacteria. In the table, the ">50" indicates that 50% inhibition of the fungus was not achieved at 50  $\mu$ g/ml which was the highest concentration tested. The abbreviation "ND" indicates that the test was not performed or that results could not be interpreted. The antimicrobial activity of MiAMP2c was also tested in the presence of 1 mM Ca<sup>2+</sup> in the test medium and the IC50 values for these tests are given in the right-hand column. As can be seen in the table, the inhibitory activity of MiAMP2c is greatly reduced (although not eliminated) in the presence of Ca<sup>2+</sup>.

Table 1

Concentrations of MiAMP2c at which 50% inhibition of growth was observed

Organism	IC50 (μg/ml)	$IC_{50} + Ca^{2+} (\mu g/ml)$
Alternaria helianthi	5-10	ND
Candida albicans	>50	>50
Ceratocystis paradoxa	20-50	>50
Cercospora nicotianae	5-10	5-10
Clavibacter michiganensis	50	>50
Chalara elegans	2-5	10-20
Fusarium oxysporum	10	20-50
Sclerotinia sclerotiorum	20-50	>50
Phytophthora cryptogea	5-10	10-25
Phytophthora parasitica nicotiana	10-20	>50

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WO 98/27805 PCT/AU97/00874 30 Verticillium dahliae 5-10 >50 Ralstonia solanacearum >50 >50 Pseudomonas syringae tabaci >50 >50 Saccharomyces cerevisiae 20-50 >50 >50 Escherichia coli >50

Table 2 shows the the antimicrobial activity of various homologues and fragments of MiAMP2c. In the table, the following abbreviations are used: Ab, *Alternaria brassicola*; Cp: *Ceratocystis paradoxa*; Foc: *Fusarium oxysporum*; Lm: *Leptosphaeria maculans*; Ss: *Sclerotinia sclerotiorum*; Vd: *Verticillium dahliae*. The ">50" indicates that concentrations higher than 50 μg/ml were not tested so that an IC50 value could not be established. A blank space indicates that the test was not performed or that results could not be interpreted.

The TcAMP1 and 2 used for the results presented in Table 2 were derived from cocoa vicilin (Examples 10 and 11). SsAMP1 and 2 show reactivity with MiAMP2c antibodies and also exhibit antimicrobial activity as seen in the table below. The versions of MiAMP2a, b and d as well as TcAMP1 and TcAMP2 tested in the bioassays all contain a His tag fusion resulting from expression in the vector pET16b. MiAMP2c pep1 and 2 are the N and C terminal regions, respectively, of MiAMP2c antimicrobial peptide as specified in Example 14 above. The concentration value listed for 'MiAMP2c pep1+2' is the concentration of each individual peptide in the mixture. It should be remembered that MiAMP2c pep1 and pep2 are both about ½ the size of MiAMP2c; comparisons of the activity of these peptides with the MiAMP2c protein should, therefore, be made on a molar basis rather than on a strict μg/ml concentration basis. Peptides were only tested in media A which did not contain added Ca<sup>2+</sup>.

Table 2

IC50 values (µg/ml) of MiAMP2 related proteins against various fungi

Peptide tested			Fungus 1	used in bioassy							
	Ab	Ср	Foc	Lm	Ss	Vd					
MiAMP2a			5-10	2.5-5	5-10						
MiAMP2b			2.5	2.5	5-10						
MiAMP2c		20-50	10		20-50	5-10					
MiAMP2d			5	2.5	5-10						
MiAMP2c pep1			100		>50						

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			31			
MiAMP2c pep2			10-20	10-20	50	10-20
MiAMP2c pep1+2			10-25		50	
TcAMP1		10	5-10	2-5	10	5-20
TcAMP2		5-10	5-10	2-5	5	5-20
SsAMP1			20-50	20-50	20-50	10-20
SsAMP2	20-50		>50	>50	>50	>50

It is worthy of note that while the TcAMP1 and 2 sequences are readily available in the public data bases, no antimicrobial activity had ever been assigned to them. These sequences were derived from much larger proteins involved in seed storage functions. The inventors have thus described a completely new activity for a small portion of the overall cocoa vicilin molecules. The activity of cotton fragments 1, 2, and 3 has been exemplified by other authors (Chung, R. P.T. *et al.* [1997] *Plant Science* 127:1-16).

#### Example 16

Construction of the plant transformation vector PCV91-MiAMP2c

The expression vector pPCV91-MiAMP2c (Figure 14) contains the full coding region of the MiAMP2c (Example 7) DNA flanked at it 5' end by the strong constitutive promoter of 35S RNA from the cauliflower mosaic virus (pCaMV35S) (Odel et al., [1985] Nature 313: 810-812) with a quadruple-repeat enhancer element (e-35S) to allow for high transcriptional activity (Kay et al. [1987] Science 236:1299-1302). The coding region of MiAMP2c DNA is flanked at its 3' end by the polyadenylation sequence of 35S RNA of the cauliflower mosaic virus (pA35S). The plasmid backbone of this vector is the plasmid pPCV91 (Walden, R. et al. [1990] Methods Mol. Cell. Biol. 1:175-194). The plasmid also contains other elements useful for plant transformation such as an ampicillin resistance gene (bla) and a hygromycin resistance gene (hph) driven by the nos promoter (pnos). These and other features allow for selection in various cloning and transformation procedures. The plasmid pPCV91-MiAMP2c was constructed as follows: A cloned fragment encoding MiAMP2c (Example 7) was digested using restriction enzymes to release the MiAMP2c gene fragment containing a synthetic leader sequence.. The binary vector pPCV91 was digested with the restriction enzyme Bam HI. Both the MiAMP2c DNA fragment containing and the binary vector were ligated using T4 DNA ligase to produce pPCV91-MiAMP2c binary vector for plant transformation (Figure 12).

Using this approach, other homologues of MiAMP2c can be expressed in plants. Not only can individual homologues be expressed, but they may be expressed in combination with other proteins as fusion proteins or as portions of larger precursor proteins. For example, it is possible to express

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the N-terminal region of MiAMP2 clone 1 (amino acids 1 to ~246) which contains a signal peptide and the hydrophilic region containing four antimicrobial segments. Transgenic plants can then be assessed to examine whether the individual fragments are being processed into the expected fragments by the processing machinery already present in the plant cells. It is also possible to express the entire MiAMP2 clone 1 (amino acids 1 to 666) and to examine the processing of the entire protein when expressed in transgenic plants. Homologous regions from other sequences can also be used in multiple combinations with, for example, ten (10) or more MiAMP2-like fragments expressed as one large fusion protein with acidic cleavage sites located as proper locations between each of the fragments. As well as linking MiAMP2 fragments together, it would also be possible to link MiAMP2 fragments to other useful proteins for expression in plants.

#### Example 17

Transgenic plants expressing MiAMP2c (or related fragments)

The disarmed *Agrobacterium tumefaciens* strain GV3101 (pMP90RK) (Koncz, Cs.[1986] *Mol. Gen. Genet.* 204:383-396) was transformed with the vector pPCV91-MiAMP2c (Example 16) using the method of Walkerpeach *et al.* (Plant Mol. Biol. Manual B1:1-19 [1994]) adapted from Van Haute *et al* (EMBO J. 2:411-417 1983]).

Tobacco transformation was carried out using leaf discs of *Nicotiana tabacum* based on the method of Horsch *et al.* (*Science* 227:1229-1231 [1985]) and co-culturing strains containing pPCV91-MiAMP2c. After co-cultivation of *Agrobacterium* and tobacco leaf disks, transgenic plants (transformed with pPCV91-MiAMP2c) were regenerated on media containing 50 µg/ml hygromycin and 500 µg/ml Cefotaxime. These transgenic plants were analysed for expression of the newly-introduced genes using standard western blotting techniques (Figure 15). Figure 15 shows a western blot of extracts from transgenic tobacco carrying the construct for MiAMP2c from example 16. Lane 1 contains pure MiAMP2c as a standard, lanes 2 and 3 contain extracts from transgenic plants carrying the pPCV91-MiAMP2c construct. As can be see in the figure, faint bands are present at approximately the correct molecular weight, indicating that the transgenic plants appear to be expressing the MiAMP2c protein. Plants capable of constitutive expression of the introduced genes may be selected and self-pollinated to give seed. F1 seedlings of the transgenic plants may be further analysed.

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## Example 18

#### MiAMP2c Homologues

Every homologue of MiAMP2c that has been tested has exhibited some antimicrobial activity. This evidence indicates that other homologues will also exhibit antimicrobial activity. These homologues include fragments from 1) peanut (Burks, A.W. et al. [1995] J. Clin. Invest. 96 (4),

1715-1721), 2) maize (Belanger, F.C. and Kriz, A.L.[1991] *Genetics* 129 (3), 863-872), 3) barley (Heck, G.R. et al. [1993] *Mol. Gen. Genet.* 239 (1-2), 209-218), and 4) soybean (Sebastiani, F.L. et al. [1990] *Plant Mol. Biol.* 15 (1), 197-201). (see SEQ ID NOS: 21, 22, 24, and 25). Other sequences derived from seed storage proteins of the 7S class are also expected to yield homologues of MiAMP2 proteins.

# SEQUENCE LISTINGS

	J	(I) GENERAL INFORMATION:													
•		(i) APPLICANT:													
		(A) NAME: COOPERATIVE RESEARCH CENTRE FOR TROPICAL PLANT PATHOLOGY													
	10	(B) STREET: The University of Queensland													
		(C) CITY: St Lucia													
		(D) STATE: Queensland													
		(E) COUNTRY: Australia (F) POSTAL CODE (ZIP): 4067													
	15	(2) 2 3 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2													
		(ii) TITLE OF INVENTION: Antimicrobial Protein													
The state of the s	20	(iii) NUMBER OF SEQUENCES: 28													
ly Lad Lad		(iv) COMPUTER READABLE FORM:													
		(A) MEDIUM TYPE: Floppy disk													
		(B) COMPUTER: IBM PC compatible													
(F)	25	<pre>(C) OPERATING SYSTEM: PC-DOS/MS-DOS (D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)</pre>													
.u   .u	(=, 101 minut 2 decire in Refease #1.0, Version #1.														
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j		(C) STRANDEDNESS: single (D) TOPOLOGY: linear													
		(2) Totomost. Hindel													
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	40	(F) TISSUE TYPE: Seeds													
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	10	Glu	Glu	Glu 115	Tyr	Asn	Arg	Gln	Arg 120	Asp	Pro	Gln	Gln	Gln 125	Tyr	Glu	Gln
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	15	Thr 145	Cys	Gln	Gln	Arg	Cys 150	Glu	Arg	Arg	Tyr	Glu 155	Lys	Glu	Lys	Arg	Lys 160
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-			610					615					620				Val
	50	625					630					635					Pro 640
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	10	(ii) MOLECULE TYPE: cDNA	
	15	<ul><li>(vi) ORIGINAL SOURCE:</li><li>(A) ORGANISM: Macadamia integrifolia</li><li>(F) TISSUE TYPE: Seeds</li></ul>	
	20	<pre>(ix) FEATURE:     (A) NAME/KEY: sig_peptide     (B) LOCATION:185</pre>	
lin, and the dim. See, A	25	<pre>(x) FEATURE:     (A) NAME/KEY: mat_peptide     (B) LOCATION:861999  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:</pre>	
d.		ATGGCGATCA ATACATCAAA TTTATGTTCT CTTCTCTTC TCCTTTCACT CTTCCTTCTG	60
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		TACTGCCAAC GACGCTGCAA GGAAATATGT GAAGAAGAAG AAGAATATAA CCGACAACGT	360
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	45	CAACAAAGA GATATGAAGA GCAACAACGT GAAGACGAAG AGAAATATGA AGAGCGAATG	540
		AAGGAAGAAG ATAACAAACG CGATCCACAA CAAAGAGAGT ACGAAGACTG CCGGAGGCGC	600
		TGCGAACAAC AGGAGCCACG TCAGCAGCAC CAGTGCCAGC TAAGATGCCG AGAGCAGCAG	660
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# (2) INFORMATION FOR SEQ ID NO: 3:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 666 amino acids

(B) TYPE: amino acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

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- (vi) ORIGINAL SOURCE:
- (A) ORGANISM: Macadamia integrifolia
- (F) TISSUE TYPE: Seeds

5 (ix) FEATURE:

- (A) NAME/KEY: sig\_peptide
- (B) LOCATION:1..28
- 10 (ix) FEATURE:
  - (A) NAME/KEY: mat peptide
  - (B) LOCATION: 29..666
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

Met Ala Ile Asn Thr Ser Asn Leu Cys Ser Leu Leu Phe Leu Leu Ser

Leu Phe Leu Leu Ser Thr Thr Val Ser Leu Ala Glu Ser Glu Phe Asp
20 25 30

Arg Gln Glu Tyr Glu Glu Cys Lys Arg Gln Cys Met Gln Leu Glu Thr 35 40 45

Ser Gly Gln Met Arg Arg Cys Val Ser Gln Cys Asp Lys Arg Phe Glu 50 60

Glu Asp Ile Asp Trp Ser Lys Tyr Asp Asn Gln Asp Asp Pro Gln Thr 65 70 75 80

Asp Cys Gln Gln Cys Gln Arg Cys Arg Gln Gln Glu Ser Gly Pro 85 90 95

Arg Gln Gln Gln Tyr Cys Gln Arg Arg Cys Lys Glu Ile Cys Glu Glu 100 105 110

Glu Glu Glu Tyr Asn Arg Gln Arg Asp Pro Gln Gln Gln Tyr Glu Gln
115 120 125

Cys Gln Glu Arg Cys Gln Arg His Glu Thr Glu Pro Arg His Met Gln
130 135 140

Thr Cys Gln Gln Arg Cys Glu Arg Arg Tyr Glu Lys Glu Lys Arg Lys 145 150 155 160

Gln Gln Lys Arg Tyr Glu Glu Gln Gln Arg Glu Asp Glu Glu Lys Tyr 165 170 175

Glu Glu Arg Met Lys Glu Glu Asp Asn Lys Arg Asp Pro Gln Gln Arg 180 185 190

Glu Tyr Glu Asp Cys Arg Arg Cys Glu Gln Gln Glu Pro Arg Gln
195 200 205

Gln Tyr Gln Cys Gln Arg Arg Cys Arg Glu Gln Gln Arg Gln His Gly
210 215 220

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		Arg 225	Gly	Gly	Asp	Leu	Ile 230	Asn	Pro	Gln	Arg	Gly 235	Gly	Ser	Gly	Arg	Tyr 240
	5	Glu	Glu	Gly	Glu	Glu 245	Lys	Gln	Ser	Asp	Asn 250	Pro	Tyr	Tyr	Phe	Asp 255	Glu
•	10	Arg	Ser	Leu	Ser 260	Thr	Arg	Phe	Arg	Thr 265	Glu	Glu	Gly	His	11e 270	Ser	Val
e .	10	Leu	Glu	Asn 275	Phe	Tyr	Gly	Arg	Ser 280	Lys	Leu	Leu	Arg	Ala 285	Leu	Lys	Asn
	15	Tyr	Arg 290	Leu	Val	Leu	Leu	Glu 295	Ala	Asn	Pro	Asn	Ala 300	Phe	Val	Leu	Pro
		Thr 305	His	Leu	Asp	Ala	Asp 310	Ala	Ile	Leu	Leu	Val 315	Thr	Gly	Gly	Arg	Gly 320
	20	Ala	Leu	Lys	Met	Ile 325	His	Arg	Asp	Asn	Arg 330	Glu	Ser	Tyr	Asn	Leu 335	Glu
	25	Cys	Gly	Asp	Val 340	Ile	Arg	Ile	Pro	Ala 345	Gly	Thr	Thr	Phe	Tyr 350	Leu	Ile
The first and first the first and	23	Asn	Arg	Asp 355	Asn	Asn	Glu	Arg	Leu 360	His	Ile	Ala	Lys	Phe 365	Leu	Gln	Thr
	30	Ile	Ser 370	Thr	Pro	Gly	Gln	Tyr 375	Lys	Glu	Phe	Phe	Pro 380	Ala	Gly	Gly	Gln
		Asn 385	Pro	Glu	Pro	Tyr	Leu 390	Ser	Thr	Phe	Ser	Lys 395	Glu	Ile	Leu	Glu	Ala 400
	35	Ala	Leu	Asn	Thr	Gln 405	Ala	Glu	Arg	Leu	Arg 410	Gly	Val	Leu	Gly	Gln 415	Gln
	40	Arg	Glu	Gly	Val 420	Ile	Ile	Ser	Ala	Ser 425	Gln	Glu	Gln	Ile	Arg 430	Glu	Leu
	10	Thr	Arg	Asp 435	Asp	Ser	Glu	Ser	Arg 440	Arg	Trp	His	Ile	Arg 445	Arg	Gly	Gly
	45	Glu	Ser 450	Ser	Arg	Gly	Pro	Tyr 455	Asn	Leu	Phe	Asn	Lys 460	Arg	Pro	Leu	Tyr
		Ser 465	Asn	Lys	Tyr	Gly	Gln 470	Ala	Tyr	Glu	Val	Lys 475	Pro	Glu	Asp	Туг	Arg 480
٠	50	Gln	Leu	Gln	Asp	Met 485		Val	Ser	Val	Phe 490	Ile	Ala	Asn	Ile	Thr 495	Gln
•	55	Gly	Ser	Met	Met 500	Gly	Pro	Phe	Phe	Asn 505	Thr	Arg	Ser	Thr	Lys 510	Val	Val
	33	Val	Val	Ala	Ser	Gly	Glu	Ala	Asp	Va1	Glu	Met	Ala	Cys	Pro	His	Leu

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515 520 525 Ser Gly Arg His Gly Gly Arg Gly Gly Lys Arg His Glu Glu Glu 5 Glu Asp Val His Tyr Glu Gln Val Lys Ala Arg Leu Ser Lys Arg Glu 550 555 Ala Ile Val Val Pro Val Gly His Pro Val Val Phe Val Ser Ser Gly 10 Asn Glu Asn Leu Leu Phe Ala Phe Gly Ile Asn Ala Gln Asn Asn 585 15 His Glu Asn Phe Leu Ala Gly Arg Glu Arg Asn Val Leu Gln Gln Ile Glu Pro Gln Ala Met Glu Leu Ala Phe Ala Ala Pro Arg Lys Glu Val 615 20 Glu Glu Leu Phe Asn Ser Gln Asp Glu Ser Ile Phe Phe Pro Gly Pro 630 635 Arg Gln His Gln Gln Ser Ser Arg Ser Thr Lys Gln Gln Fro 25 645 650 Leu Val Ser Ile Leu Asp Phe Val Gly Phe 660 30 (2) INFORMATION FOR SEQ ID NO: 4: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 2171 base pairs (B) TYPE: nucleic acid 35 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: cDNA 40 (vi) ORIGINAL SOURCE: (A) ORGANISM: Macadamia integrifolia (F) TISSUE TYPE: Seeds 45 (ix) FEATURE: (A) NAME/KEY: sig peptide (B) LOCATION:1..86 50 (ix) FEATURE: (A) NAME/KEY: mat\_peptide (B) LOCATION:87..1999 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4: 55

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TCAACGACAG TGTCTCTTGC TGAAAGTGAA TTTGACAGGC AGGAATATGA GGAGTGCAAA 120 CGGCAATGCA TGCAGTTGGA GACATCAGGC CAGATGCGTC GGTGTGTGAG TCAGTGCGAT 180 5 AAGAGATTTG AAGAGGATAT AGATTGGTCT AAGTATGATA ACCAAGACGA TCCTCAGACG 240 GATTGCCAAC AATGCCAGAG GCGATGCAGG CAGCAGGAGA GTGGCCCACG TCAGCAACAA 300 10 TACTGCCAAC GACGCTGCAA GGAAATATGT GAAGAAGAAG AAGAATATAA CCGACAACGT 360 GATCCACAGC AGCAATACGA GCAATGTCAG GAGCGCTGCC AACGGCACGA GACAGAGCCA 420 CGTCACATGC AAACATGTCA ACAACGCTGC GAGAGGAGAT ATGAAAAGGA GAAACGTAAG 480 15 CAACAAAGA GATATGAAGA GCAACAACGT GAAGACGAAG AGAAATATGA AGAGCGAATG 540 AAGGAAGAAG ATAACAAACG CGATCCACAA CAAAGAGAGT ACGAAGACTG CCGGAGGCGC 600 20 TGCGAACAAC AGGAGCCACG TCAGCAGTAC CAGTGCCAGC GAAGATGCCG AGAGCAGCAG 660 AGGCAACACG GCCGAGGTGG TGATTTGATT AACCCTCAGA GGGGAGGCAG CGGCAGATAC 720 GAGGAGGAG AAGAGAAGCA AAGCGACAAC CCCTACTACT TCGACGAACG AAGCTTAAGT 780 25 ACAAGGTTCA GGACCGAGGA AGGCCACATC TCAGTTCTGG AGAACTTCTA TGGTAGATCC 840 ı miş AAGCTTCTAC GCGCACTAAA AAACTATCGC TTGGTGCTCC TCGAGGCTAA CCCCAACGCC 900 TTCGTGCTCC CTACCCACTT GGACGCAGAT GCCATTCTCT TGGTCACCGG AGGGAGAGGA 960 ľ. GCCCTCAAAA TGATCCACCG TGACAACAGA GAATCCTACA ACCTCGAGTG TGGAGACGTA 1020 ATCAGAATCC CAGCTGGAAC CACATTCTAC TTAATCAACC GAGACAACAA CGAGAGGCTC 1080 35 CACATAGCCA AGTTCTTACA GACCATATCC ACTCCTGGCC AATACAAGGA ATTCTTCCCA 1140 GCTGGAGGCC AAAACCCAGA GCCGTACCTC AGTACCTTCA GCAAAGAGAT TCTCGAGGCT 1200 40 GCGCTCAACA CACAAGCAGA GAGGCTGCGT GGGGTGCTTG GACAGCAAAG GGAGGGAGTG 1260 ATAATTAGTG CGTCACAGGA GCAGATCAGG GAGTTGACTC GAGATGACTC AGAGTCACGA 1320 CGCTGGCATA TAAGGAGAG TGGTGAATCA AGCAGGGGAC CTTACAATCT GTTCAACAAA 1380 45 AGGCCACTGT ACTCCAACAA ATACGGTCAA GCCTACGAAG TCAAACCTGA GGACTACAGG 1440 CAACTCCAAG ACATGGACGT ATCGGTTTTC ATAGCCAACA TCACCCAGGG ATCCATGATG 1500 GGTCCCTTCT TCAACACTAG GTCTACAAAG GTGGTAGTGG TGGCTAGTGG AGAGGCAGAT 50 1560 GTGGAAATGG CATGCCCTCA CTTGTCGGGA AGACACGGCG GCCGCCGTGG AGGGAAAAGG 1620 CATGAGGAGG AAGAGGATGT GCACTATGAG CAGGTTAAAG CACGTTTGTC GAAGAGAGAG 1680 55 GCCATTGTTG TTCCGGTAGG TCATCCCGTC GTCTTCGTTT CATCCGGAAA CGAGAACCTG 1740

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	CTGCTTTTTG CATTTGGAAT CAATGCCCAA AACAACCACG AGAACTTCCT CGCGGGGAGA	1800
5	GAGAGGAACG TGCTGCAGCA GATAGAGCCA CAGGCAATGG AGCTAGCGTT TGCCGCTCCA	1860
3	AGGAAAGAGG TAGAAGAGTT ATTTAACAGC CAGGACGAGT CTATCTTCTT TCCTGGGCCC	1920
	AGGCAGCACC AGCAACAGTC TTCCCGCTCC ACCAAGCAAC AACAGCCTCT CGTCTCCATT	1980
10	CTGGACTTCG TTGGCTTCTA AAGTTCTACA AAAAAGAGTG TGTTATGTAG TATAGGTTAG	2040
	TAGCTCCTAG CTCGGTGTAT GCGAGTGGTA AGAGACCAAG ACGCTAAATC CCTAAGTAAC	2100
15	TAACCTGGCG AGCTTGCGTG TATGCAAATA AAGAGGAACA GCTTTCCAAC TTTAAAAAAA	2160
15	AAAAAAAAA A	2171
20	(2) INFORMATION FOR SEQ ID NO: 5:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 625 amino acids	
25	(B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: protein	
	(vi) ORIGINAL SOURCE:	
30	(A) ORGANISM: Macadamia integrifolia (F) TISSUE TYPE: Seeds	
	<pre>(ix) FEATURE:   (A) NAME/KEY: partial mat_peptide</pre>	
35	(B) LOCATION:1625	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:	
40	Gln Cys Met Gln Leu Glu Thr Ser Gly Gln Met Arg Arg Cys Val Ser 1 5 10 15	
	Gln Cys Asp Lys Arg Phe Glu Glu Asp Ile Asp Trp Ser Lys Tyr Asp 20 25 30	
45	Asn Gln Glu Asp Pro Gln Thr Glu Cys Gln Gln Cys Gln Arg Arg Cys 35 40 45	
50	Arg Gln Gln Glu Ser Asp Pro Arg Gln Gln Gln Tyr Cys Gln Arg Arg 50 55 60	
50	Cys Lys Glu Ile Cys Glu Glu Glu Glu Glu Tyr Asn Arg Gln Arg Asp 65 70 75 80	
55	Pro Gln Gln Gln Tyr Glu Gln Cys Gln Lys Arg Cys Gln Arg Arg Glu 85 90 95	

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		Thr	Glu	Pro	Arg 100	His	Met	Gln	Ile	Cys 105	Gln	Gln	Arg	Cys	Glu 110	Arg	Arg
	5	Tyr	Glu	Lys 115	Glu	Lys	Arg	Lys	Gln 120	Gln	Lys	Arg	Tyr	Glu 125	Glu	Gln	Gln
*		Arg	Glu 130	Asp	Glu	Glu	Lys	Tyr 135	Glu	Glu	Arg	Met	Lys 140	Glu	Gly	Asp	Asn
•	10	Lys 145	Arg	Asp	Pro	Gln	Gln 150	Arg	Glu	Tyr	Glu	Asp 155	Cys	Arg	Arg	His	Cys 160
	15	Glu 165	Gln	Gln	Glu	Pro 170	Arg	Leu	Gln	Tyr	Gln 175	Cys	Gln	Arg	Arg	Cys 180	Gln
	13	Glu	Gln	Gln	Arg 185	Gln	His	Gly	Arg	Gly 190	Gly	Asp	Leu	Met	Asn 195	Pro	Gln
le me	20	Arg	Gly	Gly 200	Ser	Gly	Arg	Tyr	Glu 205	Glu	Gly	Glu	Glu	Lys 210	Gln	Ser	Asp
		Asn	Pro 215	Tyr	Tyr	Phe	Asp	Glu 220	Arg	Ser	Leu	Ser	Thr 225	Arg	Phe	Arg	Thr
The state of the s	25	Glu 230	Glu	Gly	His	Ile	Ser 235	Val	Leu	Glu	Asn	Phe 240	Tyr	Gly	Arg	Ser	Lys 245
	30	Leu	Leu	Arg	Ala	Leu 250	Lys	Asn	Tyr	Arg	Leu 255	Val	Leu	Leu	Glu	Ala 260	Asn
ry  -		Pro	Asn	Ala	Phe 265	Val	Leu	Pro	Thr	His 270	Leu	Asp	Ala	Asp	Ala 275	Ile	Leu
S	35	Leu	Val	Ile 280	Gly	Gly	Arg	Gly	Ala 285	Leu	Lys	Met	Ile	His 290	Arg	Asp	Asn
		Arg	Glu 295	Ser	Tyr	Asn	Leu	Glu 300	Cys	Gly	Asp	Val	Ile 305	Arg	Ile	Pro	Ala
	40	Gly 310	Thr	Thr	Phe	Tyr	Leu 315	Ile	Asn	Arg	Asp	Asn 320	Asn	Glu	Arg	Leu	His 325
	45	Ile	Ala	Lys	Phe	Leu 330	Gln	Thr	Ile	Ser	Thr 335	Pro	Gly	Gln	Tyr	Lys 340	Glu
		Phe	Phe	Pro	Ala 345	Gly	Gly	Gln	Asn	Pro 350	Glu	Pro	Tyr	Leu	Ser 355	Thr	Phe
•	50	Ser	Lys	Glu 360	Ile	Leu	Glu	Ala	Ala 365	Leu	Asn	Thr	Gln	Thr 370	Glu	Arg	Leu
•		Arg	Gly 375	Val	Leu	Gly	Gln	Gln 380	Arg	Glu	Gly	Val	Ile 385	Ile	Arg	Ala	Ser
	55	Gln 390		Gln	Ile	Arg	Glu 395		Thr	Arg	Asp	Asp 400	Ser	Glu	Ser	Arg	Arg 405

			Trp	His	Ile	Arg	Arg 410	Gly	Gly	Glu	Ser	Ser 415	Arg	Gly	Pro	Tyr	Asn 420	Leu
	5		Phe	Asn	Lys	Arg 425	Pro	Leu	Tyr	Ser	Asn 430	Lys	Tyr	Gly	Gln	Ala 435	Tyr	Glu
•	10		Val	Lys	Pro 440	Glu	Asp	Tyr	Arg	Gln 445	Leu	Gln	Asp	Met	Asp 450	Val	Ser	Val
<b>=</b>	10		Phe	Ile 455	Ala	Asn	Ile	Thr	Gln 460	Gly	Ser	Met	Met	Gly 470	Pro	Phe	Phe	Asn
	15		Thr 480	Arg	Ser	Thr	Lys	Val 485	Val	Val	Val	Ala	Ser 490	Gly	Glu	Ala	Asp	Val 500
			Glu	Met	Ala	Cys	Pro 505	His	Leu	Ser	Gly	Arg 510	His	Gly	Gly	Arg	Gly 515	Gly
	20		Gly	Lys	Arg	His 520	Glu	Glu	Glu	Glu	Glu 525	Val	His	Tyr	Glu	Gln 530	Val	Arg
	25		Ala	Arg	Leu 535	Ser	Lys	Arg	Glu	Ala 540	Ile	Val	Val	Leu	Ala 545	Gly	His	Pro
in Li	23		Val	Val 550	Phe	Val	Ser	Ser	Gly 555	Asn	Glu	Asn	Leu	Leu 560	Leu	Phe	Ala	Phe
i C	30		Gly 565	Ile	Asn	Ala	Gln	Asn 570	Asn	His	Glu	Asn	Phe 575	Leu	Ala	Gly	Arg	Glu 580
iu iu j			Arg	Asn	Val	Leu	Gln 585	Gln	Ile	Glu	Pro	Gln 590	Ala	Met	Glu	Leu	Ala 595	Phe
	35		Ala	Ala	Ser	Arg 600	Lys	Glu	Va1	Glu	Glu 605	Leu	Phe	Asn	Ser	Gln 610	Asp	Glu
	40		Ser	Ile	Phe 615	Phe	Pro	Gly	Pro	Arg 620	Gln	His	Gln	Gln	Gln 625	Ser	Pro	Arg
	70		Ser Phe	Thr 630	Lys	Gln	Gln	Gln	Pro 635		Val	Ser	Ile	Leu 640	Asp	Phe	Val	Gly
	45	(2)	INFO	RMAT	'ION	FOR	SEQ	ID N	O: 6	:								
			(i)	(A)	) LEI	E CH. NGTH PE: 1	: 214	40 b	ase j	pair	S							
•	50			(C)		RANDI POLO			_	le								
•			(ii	) MO	LECU	LE T	YPE:	cDN	A									
	55		(vi	) OR	.IGIN	AL S	OURC	E:										

(A) ORGANISM: Macadamia integrifolia

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(F) TISSUE TYPE: Seeds

## (x) FEATURE:

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(A) NAME/KEY: partial mat\_peptide

(B) LOCATION:1..1875

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

•	10	CAATGCATGC	AGTTAGAGAC	ATCAGGCCAG	ATGCGTCGGT	GTGTGAGTCA	GTGCGATAAG	60
2	10	AGATTTGAAG	AGGATATAGA	TTGGTCTAAG	TATGATAACC	AAGAGGATCC	TCAGACGGAA	120
		TGCCAACAAT	GCCAGAGGCG	ATGCAGGCAG	CAGGAGAGTG	ACCCACGTCA	GCAACAATAC	180
	15	TGCCAACGAC	GCTGCAAGGA	AATATGTGAA	GAAGAAGAAG	AATATAACCG	ACAACGTGAT	240
		CCACAGCAGC	AATACGAGCA	ATGTCAGAAG	CGCTGCCAAC	GGCGCGAGAC	AGAGCCACGT	300
	20	CACATGCAAA	TATGTCAACA	ACGCTGCGAG	AGGAGATATG	AAAAGGAGAA	ACGTAAGCAA	360
43 L	20	CAAAAGAGAT	ATGAAGAGCA	ACAACGTGAA	GACGAAGAGA	AATATGAAGA	GCGAATGAAG	420
		GAAGGAGATA	ACAAACGCGA	TCCACAACAA	AGAGAGTACG	AAGACTGCCG	GCGGCACTGC	480
	25	GAACAACAGG	AGCCACGTCT	GCAGTACCAG	TGCCAGCGAA	GATGCCAAGA	GCAGCAGAGG	540
4		CAACACGGCC	GAGGTGGCGA	TTTGATGAAC	CCTCAGAGGG	GAGGCAGCGG	CAGATACGAG	600
H	20	GAGGGAGAAG	AGAAGCAAAG	CGACAACCCC	TACTACTTCG	ACGAACGAAG	CTTAAGTACA	660
	30	AGGTTCAGGA	CCGAGGAAGG	CCACATCTCA	GTTCTGGAGA	ACTTCTATGG	TAGATCCAAG	720
A Comment		CTTCTACGCG	CACTAAAAAA	CTATCGCTTG	GTGCTCCTCG	AGGCTAACCC	CAACGCCTTC	780
	35	GTGCTCCCTA	CCCACTTGGA	TGCAGATGCC	ATTCTCTTGG	TCATCGGAGG	GAGAGGAGCC	840
		CTCAAAATGA	TCCACCGTGA	CAACAGAGAA	TCCTACAACC	TCGAGTGTGG	AGACGTAATC	900
	4.0	AGAATCCCAG	CTGGAACCAC	ATTCTACTTA	ATCAACCGAG	ACAACAACGA	GAGGCTCCAC	960
	40	ATAGCCAAGT	TCTTACAGAC	CATATCCACT	CCTGGCCAAT	ACAAGGAATT	CTTCCCAGCT	1020
		GGAGGCCAAA	ACCCAGAGCC	GTACCTCAGT	ACCTTCAGCA	AAGAGATTCT	CGAGGCTGCG	1080
	45	CTCAACACAC	AAACAGAGAG	GCTGCGTGGG	GTGCTTGGAC	AGCAAAGGGA	GGGAGTGATA	1140
		ATTAGGGCGT	CACAGGAGCA	GATCAGGGAG	TTGACTCGAG	ATGACTCAGA	GTCACGACGC	1200
		TGGCATATAA	GGAGAGGTGG	TGAATCAAGC	AGGGGACCTT	ACAATCTGTT	CAACAAAAGG	1260
•	50	CCACTGTACT	CCAACAAATA	CGGTCAAGCC	TACGAAGTCA	AACCTGAGGA	CTACAGGCAA	1320
_		CTCCAAGACA	TGGACGTATC	AGTTTTCATA	GCCAACATCA	CCCAGGGATC	CATGATGGGT	1380
•	55	CCCTTCTTCA	ACACTAGGTC	TACAAAGGTG	GTAGTGGTGG	CTAGTGGAGA	GGCAGATGTG	1440

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	GAAATGGCAT GCCCTCACTT GTCGGGAAGA CACGGCGGCC GCGGTGGAGG GAAAAGGCAT	1500
	GAGGAGGAG AGGAGGTGCA CTATGAGCAG GTTAGAGCAC GTTTGTCGAA GAGAGAGGCC	1560
5	ATTGTTGTTC TGGCAGGTCA TCCCGTCGTC TTCGTTTCAT CCGGAAACGA AAACCTGCTG	1620
	CTTTTTGCAT TTGGAATCAA TGCCCAAAAC AACCACGAGA ACTTCCTCGC GGGGAGAGAG	1680
10	AGGAACGTGC TGCAGCAGAT AGAGCCACAG GCAATGGAGC TAGCGTTTGC CGCTTCAAGG	1740
10	AAAGAGGTAG AAGAGTTATT TAACAGCCAG GACGAGTCTA TCTTCTTTCC TGGGCCCAGG	1800
	CAGCACCAGC AACAGTCGCC CCGCTCCACC AAGCAACAAC AGCCTCTCGT CTCCATTCTG	1860
. 15	GACTTCGTTG GCTTCTAAAG TTCTACAAAA AAGAGTGTGT TATGTAGTAT AGGTTAGTAG	1920
	CTCCTAGCTC GGTGTATGAG AGTGGTAAGA GACTAAGACG CTAAATCCCT AAGTAACTAA	1980
20	CCTGGCGAGC TTGCGTGTAT GCAAATAAAG AGGAACAGCT TTCCAACTTT AGAAAGCTCT	2040
20	TTTTTTTTT TTTTTTCTTT CTTTTCTTA AGAAATAAAC GAACGTAGAT TGCGGCTCAA	2100
•	AAAAAAAA AAAAAAAA AAAAAAAAA AAAAAAAAA	2140
25	(2) INFORMATION FOR SEQ ID NO: 7:	
30	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 525 amino acids</li> <li>(B) TYPE: amino acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
35	<ul><li>(ii) MOLECULE TYPE: protein</li><li>(vi) ORIGINAL SOURCE:</li><li>(A) ORGANISM: Theobroma cacao</li><li>(F) TISSUE TYPE: Seeds</li></ul>	
40	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:	
	Met Val Ile Ser Lys Ser Pro Phe Ile Val Leu Ile Phe Ser Leu Leu 1 5 10 15	
45	Leu Ser Phe Ala Leu Leu Cys Ser Gly Val Ser Ala Tyr Gly Arg Lys 20 25 30	
50	Gln Tyr Glu Arg Asp Pro Arg Gln Gln Tyr Glu Gln Cys Gln Arg Arg 35 40 45	
50	Cys Glu Ser Glu Ala Thr Glu Glu Arg Glu Gln Glu Gln Cys Glu Gln 50 55 60	
55	Arg Cys Glu Arg Glu Tyr Lys Glu Gln Gln Arg Gln Gln Glu Glu 65 70 75 80	

		Leu	Gln	Arg	Gln	Tyr 85	Gln	Gln	Cys	Gln	Gly 90	Arg	Cys	Gln	Glu	Gln 95	Gln
	5	Gln	Gly	Gln	Arg 100	Glu	Gln	Gln	Gln	Cys 105	Gln	Arg	Lys	Cys	Trp 110	Glu	Gln
ų.		Tyr	Lys	Glu 115	Gln	Glu	Arg	Gly	Glu 120	His	Glu	Asn	Tyr	His 125	Asn	His	Lys
¥,	10	Lys	Asn 130	Arg	Ser	Glu	Glu	Glu 135	Glu	Gly	Gln	Gln	Arg 140	Asn	Asn	Pro	Tyr
	15	Tyr 145	Phe	Pro	Lys	Arg	Arg 150	Ser	Phe	Gln	Thr	Arg 155	Phe	Arg	Asp	Glu	Glu 160
		Gly	Asn	Phe	Lys	Ile 165	Leu	Gln	Arg	Phe	Ala 170	Glu	Asn	Ser	Pro	Pro 175	Leu
	20	_	_	Ile	180		_	_		185					190		
to decrease the tens does to the				Ile 195					200					205			
	25		210	Gly				215					220				
	30	225		Asn			230	_				235				-	240
				Tyr		245			_		250		-			255	
	35			Ala	260					265		-			270		
	40			Gly 275 Leu					280				_	285			-
	40		290	Glu				295					300				
	45	305		Arg			310	-				315		-			320
				Ser		325					330					335	
ŧ	50			Gln	340					345					350		
*	55			355 Pro					360					365			
			370			•		375			_		380	L			

Ser Ala Phe Lys Leu Asn Gln Gly Ala Ile Phe Val Pro His Tyr Asn 385 390 5 Ser Lys Ala Thr Phe Val Val Phe Val Thr Asp Gly Tyr Gly Tyr Ala 410 Gln Met Ala Cys Pro His Leu Ser Arg Gln Ser Gln Gly Ser Gln Ser 10 Gly Arg Gln Asp Arg Glu Glu Glu Glu Glu Glu Glu Glu Glu Thr 435 440 Phe Gly Glu Phe Gln Gln Val Lys Ala Pro Leu Ser Pro Gly Asp Val 15 455 Phe Val Ala Pro Ala Gly His Ala Val Thr Phe Phe Ala Ser Lys Asp 465 470 475 20 Gln Pro Leu Asn Ala Val Ala Phe Gly Leu Asn Ala Gln Asn Asn Gln Arg Ile Phe Leu Ala Gly Arg Pro Phe Phe Leu Asn His Lys Gln Asn 505 25 Thr Asn Val Ile Lys Phe Thr Val Lys Ala Ser Ala Tyr 515 520 30 (2) INFORMATION FOR SEQ ID NO: 8: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 590 amino acids (B) TYPE: amino acid 35 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein 40 (vi) ORIGINAL SOURCE: (A) ORGANISM: Gossypium hirsutum (F) TISSUE TYPE: Seeds (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8: 45 Met Val Arg Asn Lys Ser Ala Cys Val Val Leu Leu Phe Ser Leu Phe 10 Leu Ser Phe Gly Leu Leu Cys Ser Ala Lys Asp Phe Pro Gly Arg Arg 50 20 25 Gly Asp Asp Pro Pro Lys Arg Tyr Glu Asp Cys Arg Arg Arg Cys 55 Glu Trp Asp Thr Arg Gly Gln Lys Glu Gln Gln Gln Cys Glu Glu Ser

		Cys 65	Lys	Ser	Gln	Tyr	Gly 70	Glu	Lys	Asp	Gln	Gln 75	Gln	Arg	His	Arg	Pro 80
	5	Glu	Asp	Pro	Gln	Arg 85	Arg	Tyr	Glu	Glu	Cys 90	Gln	Gln	Glu	Cys	Arg 95	Gln
•	10	Gln	Glu	Glu	Arg 100	Gln	Gln	Pro	Gln	Cys 105	Gln	Gln	Arg	Cys	Leu 110	Lys	Arg
*		Phe	Glu	Gln 115	Glu	Gln	Gln	Gln	Ser 120	Gln	Arg	Gln	Phe	Gln 125	Glu	Cys	Gln
	15	Gln	His 130	Cys	His	Gln	Gln	Glu 135	Gln	Arg	Pro	Glu	Lys 140	Lys	Gln	Gln	Cys
		Val 145	Arg	Glu	Cys	Arg	Glu 150	Lys	Tyr	Gln	Glu	Asn 155	Pro	Trp	Arg	Gly	Glu 160
A STATE OF THE STA	20	Arg	Glu	Glu	Glu	Ala 165	Glu	Glu	Glu	Glu	Thr 170	Glu	Glu	Gly	Glu	Gln 175	Glu
	25	Gln	Ser	His	Asn 180	Pro	Phe	His	Phe	His 185	Arg	Arg	Ser	Phe	Gln 190	Ser	Arg
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		Phe	Arg	Glu 195	Glu	His	Gly	Asn	Phe 200	Arg	Val	Leu	Gln	Arg 205	Phe	Ala	Ser
	30	Arg	His 210	Pro	Ile	Leu	Arg	Gly 215	Ile	Asn	Glu	Phe	Arg 220	Leu	Ser	Ile	Leu
		225	Ala				230					235					240
1 100	35		Ile			245					250					255	
	40		Glu		260					265					270		
			Pro	275					280					285			
	45		Leu 290					295					300			_	
		305	Glu				310					315					320
*	50		Ala			325					330					335	
4	55		Gln		340					345					350		
		Gln	Gly	Gln	Gly	Met	Phe	Arg	Lys	Ala	Ser	Gln	Glu	Gln	Ile	Arg	Ala

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355 ` 360 Leu Ser Gln Glu Ala Thr Ser Pro Arg Glu Lys Ser Gly Glu Arg Phe 375 5 Ala Phe Asn Leu Leu Ser Gln Thr Pro Arg Tyr Ser Asn Gln Asn Gly 385 390 395 Arg Phe Phe Glu Ala Cys Pro Pro Glu Phe Arg Gln Leu Arg Asp Ile 10 Asn Val Thr Val Ser Ala Leu Gln Leu Asn Gln Gly Ser Ile Phe Val 420 425 15 Pro His Tyr Asn Ser Lys Ala Thr Phe Val Ile Leu Val Thr Glu Gly 440 Asn Gly Tyr Ala Glu Met Val Ser Pro His Leu Pro Arg Gln Ser Ser 455 20 465 470 475 Glu Glu Arg Arg Ser Gly Gln Tyr Arg Lys Ile Arg Ser Arg Leu Ser 25 Arg Gly Asp Ile Phe Val Val Pro Ala Asn Phe Pro Val Thr Phe Val 505 Ala Ser Gln Asn Gln Asn Leu Arg Met Thr Gly Phe Gly Leu Tyr Asn 30 515 Gln Asn Ile Asn Pro Asp His Asn Gln Arg Ile Phe Val Ala Gly Lys 535 35 Ile Asn His Val Arg Gln Trp Asp Ser Gln Ala Lys Glu Leu Ala Phe 545 550 Gly Val Ser Ser Arg Leu Val Asp Glu Ile Phe Asn Ser Asn Pro Gln 40 Glu Ser Tyr Phe Val Ser Arg Gln Arg Gln Arg Ala Ser Glu 585 45 (2) INFORMATION FOR SEQ ID NO: 9: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 22 amino acids (B) TYPE: amino acid 50 (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:

Arg Gln Arg Asp Pro Gln Gln Gln Ala Glu Gln Ala Gln Lys Arg Ala 10 Gln Arg Arg Glu Thr Glu 5 20 (2) INFORMATION FOR SEQ ID NO: 10: 10 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 15 (ii) MOLECULE TYPE: protein (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10: 20 Pro Arg His Met Gln Ile Ala Gln Gln Arg Ala Glu Arg Arg Ala Glu The state of the s 10 Lys Glu Lys Arg Lys Gln Gln Lys Arg 20 25 (2) INFORMATION FOR SEQ ID NO: 11: (i) SEQUENCE CHARACTERISTICS: 30 (A) LENGTH: 30 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear J 35 (ii) MOLECULE TYPE: protein (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11: Met Ala Trp Phe His Val Ser Val Cys Asn Ala Val Phe Val Val Ile 40 Ile Ile Ile Met Leu Leu Met Phe Val Pro Val Val Arg Gly 20 25 30 45 (2) INFORMATION FOR SEQ ID NO: 12: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs 50 (B) TYPE: nucleotide (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: nucleotide 55 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:

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		CAGCAGCAGT ATGAGCAGTG	20
	5	(2) INFORMATION FOR SEQ ID NO: 13:	
	10	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
		(ii) MOLECULE TYPE: DNA	
	15	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:	
		TTTTTCGTAK CKKCKTTCGC A	21
2 800 g	20	(2) INFORMATION FOR SEQ ID NO: 14:	
The state of the s	25	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 24 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
end,		(ii) MOLECULE TYPE: DNA	
1,00	30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:	
		ACACCATATG CGACAACGTG ATCC	24
	35	(2) INFORMATION FOR SEQ ID NO: 15:	
	40	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 26 base pairs</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>	
		(ii) MOLECULE TYPE: DNA	
	45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:	
		CGTTGTTTTC TCTATTCCTA GGGTTG	26
	50	(2) INFORMATION FOR SEQ ID NO: 16:	
ţ	55	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 22 amino acids</li><li>(B) TYPE: amino acid</li><li>(C) STRANDEDNESS: single</li></ul>	
		(D) TOPOLOGY: linear	

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	(ii) MOLECULE TYPE: protein	
5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:	
3	Met Gly His His His His His His His His His Ser Ser Gly His 1 5 10 15	
10	Ile Glu Gly Arg His Met 20	
	(2) INFORMATION FOR SEQ ID NO: 17:	
15	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 90 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
20	(ii) MOLECULE TYPE: DNA	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17	
25	GGGAATTCCA TATGTATGAG CGTGATCCTC GACAGCAATA CGAGCAATGC CAGAGGCGAT	60
	GCGAGTCGGA AGCGACTGAA GAAAGGGAGC	90
30	(2) INFORMATION FOR SEQ ID NO: 18	
35	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 91 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
	(ii) MOLECULE TYPE: DNA	
40	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 18:	
	GAAGCGACTG AAGAAAGGGA GCAAGAGCAG TGTGAACAAC GCTGTGAAAG GGAGTACAAG	60
45	GAGCAGCAGA GACAGCAATA GGGATCCACA C	91
	(2) INFORMATION FOR SEQ ID NO: 19	
50	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 101 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
55	(ii) MOLECULE TYPE: DNA	

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		(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 19:	
		GGGAATTCCA TATGCTTCAA AGGCAATACC AGCAATGTCA AGGGCGTTGT CAAGAGCAAC	60
	5	AACAGGGCA GAGAGAGCAG CAGCAGTGCC AGAGAAAATG C	101
		(2) INFORMATION FOR SEQ ID NO: 20	
	10	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 102 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
	13	(ii) MOLECULE TYPE: DNA	
		(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 20	
	20	GTGTGGATCC CTAGCTCCTA TTTTTTTTGT GATTATGGTA ATTCTCGTGC TCGCCTCTCT	60
		CTTGTTCCTT ATATTGCTCC CAGCATTTTC TCTGGCACTG CT	102
	25	(2) INFORMATION FOR SEQ ID NO: 21:	
	30	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 42 amino acids</li> <li>(B) TYPE: amino acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
	35	(ii) MOLECULE TYPE: protein	
•		<pre>(vi) ORIGINAL SOURCE:   (A) ORGANISM: Peanut   (F) TISSUE TYPE: Seeds</pre>	
	40	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 21:	
		Met Arg Gly Arg Val Ser Pro Leu Met Leu Leu Gly Ile Leu Val 1 5 10 15	
	45	Leu Ala Ser Val Ser Ala Thr Gln Ala Lys Ser Pro Tyr Arg Lys Thr 20 25 30	
	50	Glu Asn Pro Cys Ala Gln Arg Cys Leu Gln Ser Cys Gln Gln Glu Pro 35 40 45	
	50	Asp Asp Leu Lys Gln Lys Ala Cys Glu Ser Arg Cys Thr Lys Leu Glu 50 55 60	
	55	Tyr Asp Pro Arg Cys Val Tyr Asp Thr Gly Ala Thr Asn Gln Arg His 65 70 75 80	

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		Pro	Pro	Gly	Glu	Arg 85	Thr	Arg	Gly	Arg	Gln 90	Pro	Gly	Asp	Tyr	Asp 95	Asp
	5	Asp	Arg	Arg	Gln 100	Pro	Arg	Arg	Glu	Glu 105	Gly	Gly	Arg	Trp	Gly 110	Pro	Ala
*		Glu	Pro	Arg 115	Glu	Arg	Glu	Arg	Glu 120	Glu	Asp	Trp	Arg	Gln 125	Pro	Arg	Glu
	10	Asp	Trp 130	Arg	Arg	Pro	Ser	His 135	Gln	Gln	Pro	Arg	Lys 140	Ile	Arg	Pro	Glu
	15	Gly 145	Arg	Glu	Gly	Glu	Gln 150	Glu	Trp	Gly	Thr	Pro 155	Gly	Ser	Glu	Val	Arg 160
		Glu 165	Glu	Thr	Ser	Arg 170	Asn	Asn	Pro	Phe	Tyr 175	Phe	Pro	Ser	Arg	Arg 180	Phe
	20			Arg	185	-				190			J		195		J
The state state start start	0.5		-	Gln 200			_		205					210		J	
	25		215	Ile			_	220					225		-		
	30	230		Asp			235					240					245
				Asn	-	250			-		255			-		260	
	35			Gln	265					270					275		
	40			280 Gln					285					290			
	10		295					300					305				Asn
	45	310	-	Phe			315		_			320					325
						330					335					340	Ser
•	50	-			345		_	-		350					355		Glu
٥	55			360					365					370			Glu
			375	-10	~		70	380			<i>y</i> ~	-10	385		JLU	J. U	<b></b> u

		390	Ile	Thr	Asn	Pro	11e 395	Asn	Leu	Arg	Asp	Gly 400	Glu	Pro	Asp	Leu	Ser 405
	5	Asn	Asn	Phe	Gly	Arg 410	Leu	Phe	Glu	Val	Lys 415	Pro	Asp	Lys	Lys	Asn 420	Pro
•	10	Gln	Leu	Gln	Asp 425	Leu	Asp	Met	Met	Leu 430	Thr	Cys	Val	Glu	Ile 435	Lys	Glu
ı	•	Gly	Ala	Leu 440	Met	Leu	Pro	His	Phe 445	Asn	Ser	Lys	Ala	Met 450	Val	Ile	Val
	15	Val	Val 455	Asn	Lys	Gly	Thr	Gly 460	Asn	Leu	Glu	Leu	Val 470	Ala	Val	Arg	Lys
		Glu 480	Gln	Gln	Gln	Arg	Gly 485	Arg	Arg	Glu	Gln	Glu 490	Trp	Glu	Glu	Glu	Glu 500
	20	Glu	Asp	Glu	Glu	Glu 505	Glu	Gly	Ser	Asn	Arg 510	Glu	Val	Arg	Arg	Tyr 515	Thr
The state of the s	25	Ala	Arg	Leu	Lys 520	Glu	Gly	Asp	Val	Phe <b>52</b> 5	Ile	Met	Pro	Ala	Ala 530	His	Pro
		Val	Ala	Ile 535	Asn	Ala	Ser	Ser	Glu 540	Leu	His	Leu	Leu	Gly 545	Phe	Gly	Ile
: (1)	30	Asn	Ala 550	Glu	Asn	Asn	His	Arg 555	Ile	Phe	Leu	Ala	Gly 560	Asp	Lys	Asp	Asn
		Val 565	Ile	Asp	Gln	Ile	Glu 570	Lys	Gln	Ala	Lys	Asp 575	Leu	Ala	Phe	Pro	Gly 580
	35	Ser	Gly	Glu	Gln	Val 585	Glu	Lys	Leu	Ile	Lys 590	Asn	Gln	Arg	Glu	Ser 595	His
	40	Phe	Val	Ser	Ala 600	Arg	Pro	Gln	Ser	Gln 605	Ser	Pro	Ser	Ser	Pro 610	Glu	Lys
	40	Glu	Asp	Gln 615	Glu	Glu	Glu	Asn	Gln 620	_	Gly	Lys	Gly	Pro 625	Leu	Leu	Ser
	45	Ile	Leu 630	Lys	Ala	Phe	Asn										
		(2)	INF	'AMRC	TION	FOR	SEQ	ID 1	NO: :	22:							

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(i) SEQUENCE CHARACTERISTICS:

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- (A) LENGTH: 46 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

- - (F) TISSUE TYPE: Seeds

5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 22: Met Val Ser Ala Arg Ile Val Val Leu Leu Ala Thr Leu Leu Cys Ala 10 Ala Ala Ala Val Ala Ser Ser Trp Glu Asp Asp Asn His His His 25 Gly Gly His Lys Ser Gly Gln Cys Val Arg Arg Cys Glu Asp Arg Pro 15 Trp His Gln Arg Pro Arg Cys Leu Glu Gln Cys Arg Glu Glu Glu Arg 50 Glu Lys Arg Gln Glu Arg Ser Arg His Glu Ala Asp Asp Arg Ser Gly 20 Glu Gly Ser Ser Glu Asp Glu Arg Glu Gln Glu Lys Glu Lys Gln Lys Asp Arg Arg Pro Tyr Val Phe Asp Arg Arg Ser Phe Arg Arg Val Val sal, m m m Arg Ser Glu Gln Gly Ser Leu Arg Val Leu Arg Pro Phe Asp Glu Val 30 Ser Arg Leu Leu Arg Gly Ile Arg Asp Tyr Arg Val Ala Val Leu Glu :=; 130 ū Ala Asn Pro Arg Ser Phe Val Val Pro Ser His Thr Asp Ala His Cys 35 150 Ile Cys Tyr Val Ala Glu Gly Glu Gly Val Val Thr Thr Ile Glu Asn 165 175 40 Gly Glu Arg Arg Ser Tyr Thr Ile Lys Gln Gly His Val Phe Val Ala 185 Pro Ala Gly Ala Val Thr Tyr Leu Ala Asn Thr Asp Gly Arg Lys 45 205 Leu Val Ile Thr Lys Ile Leu His Thr Ile Ser Val Pro Gly Glu Phe 215 220 50 Gln Phe Phe Phe Gly Pro Gly Gly Arg Asn Pro Glu Ser Phe Leu Ser 240 Ser Phe Ser Lys Ser Ile Gln Arg Ala Ala Tyr Lys Thr Ser Ser Asp 250 55 Arg Leu Glu Arg Leu Phe Gly Arg His Gly Gln Asp Lys Gly Ile Ile

Val Arg Ala Thr Glu Glu Gln Thr Arg Glu Leu Arg Arg His Ala Ser Glu Gly Gly His Gly Pro His Trp Pro Leu Pro Pro Phe Gly Glu Ser Arg Gly Pro Tyr Ser Leu Leu Asp Gln Arg Pro Ser Ile Ala Asn Gln His Gly Gln Leu Tyr Glu Ala Asp Ala Arg Ser Phe His Asp Leu Ala Glu His Asp Val Ser Val Ser Phe Ala Asn Ile Thr Ala Gly Ser Met Ser Ala Pro Leu Phe Asn Thr Arg Ser Phe Lys Ile Ala Tyr Val Pro Asn Gly Lys Gly Tyr Ala Glu Ile Val Cys Pro His Arg Gln Ser Gln Gly Gly Glu Ser Glu Arg Glu Arg Asp Lys Gly Arg Arg Ser Glu Glu L. Glu Glu Glu Glu Ser Ser Glu Glu Glu Glu Glu Ala Gly Gln Gly Tyr His Thr Ile Arg Ala Arg Leu Ser Pro Gly Thr Ala Phe Val Val Pro a ===== Ala Gly His Pro Phe Val Ala Val Ala Ser Arg Asp Ser Asn Leu Gln Ile Val Cys Phe Glu Val His Ala Asp Arg Asn Glu Lys Val Phe Leu Ala Gly Ala Asp Asn Val Leu Gln Lys Leu Asp Arg Val Ala Lys Ala Leu Ser Phe Ala Ser Lys Ala Glu Glu Val Asp Glu Val Leu Gly Ser Arg Arg Glu Lys Gly Phe Leu Pro Gly Pro Glu Glu Ser Gly Gly His Glu Glu Arg Glu Gln Glu Glu Glu Arg Glu Glu Arg His Gly Gly Arg Gly Glu Arg Glu Arg His Gly Arg Glu Glu Arg Glu Lys Glu Glu Glu Arg Glu Gly Arg His Gly Gly Arg Glu Glu Arg Glu Glu Glu

Arg His Gly Arg Gly Arg Glu Glu Val Ala Glu Thr Leu Met Arg
585 590 595

Met Val Thr Ala Arg Met 5 600

(2) INFORMATION FOR SEQ ID NO: 23:

10 (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 33 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Maize
- (F) TISSUE TYPE: Seeds
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 23:

Arg Ser Gly Arg Gly Glu Cys Arg Arg Gln Cys Leu Arg Arg His Glu
25 1 5 10 15

Gly Gln Pro Trp Glu Thr Gln Glu Cys Met Arg Arg Cys Arg Arg Arg 20 25 30

30 Gly

- (2) INFORMATION FOR SEQ ID NO: 24:
- 35 (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 42 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Barley
- 45 (F) TISSUE TYPE: Seeds
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 24:

Met Ala Thr Arg Ala Lys Ala Thr Ile Pro Leu Leu Phe Leu Leu Gly
50 1 5 10 15

Thr Ser Leu Leu Phe Ala Ala Ala Val Ser Ala Ser His Asp Asp Glu 20 25 30

55 Asp Asp Arg Gly Gly His Ser Leu Gln Gln Cys Val Gln Arg Cys 35 40 45

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		Arg	Gln 50	Glu	Arg	Pro	Arg	Tyr 55	Ser	His	Ala	Arg	Cys 60	Val	Gln	Glu	Cys
	5	Arg 65	Asp	Asp	Gln	Gln	Gln 70	His	Gly	Arg	His	Glu 75	Gln	Glu	Glu	Glu	Gln 80
	10	Gly	Arg	Gly	Arg	Gly 85	Trp	His	Gly	Glu	Gly 90	Glu	Arg	Glu	Glu	Glu 95	His
	10	Gly	Arg	Gly	Arg 100	Gly	Arg	His	Gly	Glu 105	Gly	Glu	Arg	Glu	Glu 110	Glu	His
	15	Gly	Arg	Gly 115	Arg	Gly	Arg	His	Gly 120	Glu	Gly	Glu	Arg	Glu 125	Glu	Glu	Arg
		Gly	Arg 130	Gly	His	Gly	Arg	His 135	Gly	Glu	Gly	Glu	Arg 140	Glu	Glu	Glu	Arg
	20	Gly 145	Arg	Gly	Arg	Gly	Arg 150	His	Gly	Glu	Gly	Glu 155	Arg	Glu	Glu	Glu	Glu 160
	25	Gly 165	Arg	Gly	Arg	Gly 170	Arg	Arg	Gly	Glu	Gly 175	Glu	Arg	Asp	Glu	Glu 180	Gln
		Gly	Asp	Ser	Arg 185	Arg	Pro	Tyr	Val	Phe 190	Gly	Pro	Arg	Ser	Phe 195	Arg	Arg
13	30	Ile	Ile	Gln 200	Ser	Asp	His	Gly	Phe 205	Val	Arg	Ala	Leu	Arg 210	Pro	Phe	Asp
		Gln	Val 215	Ser	Arg	Leu	Leu	Arg 220	Gly	Ile	Arg	Asp	Tyr 225	Arg	Val	Ala	Ile
	35	Met 230	Glu	Val	Asn	Pro	Arg 235	Ala	Phe	Val	Val	Pro 240	Gly	Phe	Thr	Asp	Ala 245
	40	Asp	Gly	Val	Gly	Tyr 250	Val	Ala	Gln	Gly	Glu 255	Gly	Val	Leu	Thr	Val 260	Ile
	••	Glu	Asn	Gly	Glu 265	Lys	Arg	Ser	Tyr	Thr 270	Val	Lys	Glu	Gly	Asp 275	Val	Ile
	45	Val	Ala	Pro 280	Ala	Gly	Ser	Ile	Met 285	His	Leu	Ala	Asn	Thr 290	Asp	Gly	Arg
		Arg	Lys 295	Leu	Val	Ile	Ala	100	Ile	Leu	His	Thr	Ile 305	Ser	Val	Pro	Gly
•	50	Lys 310	Phe	Gln	Phe	Leu	Ser 315	Val	Lys	Pro	Leu	Leu 320	Ala	Ser	Leu	Ser	Lys 325
•	55	Arg	Val	Leu	Arg	Ala 330	Ala	Phe	Lys	Thr	Ser 335	Asp	Glu	Arg	Leu	Glu 340	Arg
		Leu	Phe	Asn	Gln	Arg	Gln	Gly	Gln	Glu	Lys	Thr	Arg	Ser	Val	Ser	Ile

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62 345 350 355 Val Arg Ala Ser Glu Glu Gln Leu Arg Glu Leu Arg Arq Glu Ala Ala 365 5 Glu Gly Gly Gln Gly His Arg Trp Pro Leu Pro Pro Phe Arg Gly Asp 380 Ser Arg Asp Thr Phe Asn Leu Leu Glu Gln Arg Pro Lys Ile Ala Asn 10 395 Arg His Gly Arg Leu Tyr Glu Ala Asp Ala Arg Ser Phe His Ala Leu 415 15 Ala Asn Gln Asp Val Arg Val Ala Val Ala Asn Ile Thr Pro Gly Ser Met Thr Ala Pro Tyr Leu Asn Thr Gln Ser Phe Lys Leu Ala Val Val 20 Leu Glu Gly Glu Gly Glu Val Gln Ile Val Cys Pro His Leu Gly Arg Glu Ser Glu Ser Glu Arg Glu His Gly Lys Gly Arg Arg Arg Glu Glu 25 485 490 Glu Glu Asp Asp Gln Arg Gln Gln Arg Arg Arg Gly Ser Glu Ser Glu 510 30 Ser Glu Glu Glu Glu Gln Gln Arg Tyr Glu Thr Val Arg Ala Arg Val Ser Arg Gly Ser Ala Phe Val Val Pro Pro Gly His Pro Val Val 540 35 Glu Ile Ser Ser Ser Gln Gly Ser Ser Asn Leu Gln Val Val Cys Phe 555 Glu Ile Asn Ala Glu Arg Asn Glu Arg Val Trp Leu Ala Gly Arg Asn 40 570 Asn Val Ile Gly Lys Leu Gly Ser Pro Ala Gln Glu Leu Thr Phe Gly 585 590 45 Arg Pro Ala Arg Glu Val Gln Glu Val Phe Arg Ala Gln Asp Gln Asp Glu Gly Phe Val Ala Gly Pro Glu Gln Gln Ser Arg Glu Gln Glu Gln 620 50 Glu Gln Glu Arg His Arg Arg Gly Asp Arg Gly Arg Gly Asp Glu Ala Val Glu Thr Phe Leu Arg Met Ala Thr Gly Ala Ile 55 645 650 655

	(2)	INF	ORMA	TION	FOR	SEQ	ID	NO:	25:							
5		(i	( <i>I</i> (I	A) LE 3) T C) S	CE C ENGTI YPE: TRANI OPOLO	H: 59 amir DEDNI	am: no ac ESS:	ino a cid sing	acids	3						
10		(i	i) M	OLECI	ULE '	TYPE	: pro	otei	n							
15			(F	A) OF	RGANI	SM: TYP	Soyk E: S	seeds	3							
		(x:	i) SI	EQUE	NCE I	DESCI	RIPT:	ON:	SEQ	ID 1	10: 2	25:				
20	Met 1	Met	Arg	Ala	Arg 5	Phe	Pro	Leu	Leu	Leu 10	Leu	Gly	Leu	Val	Phe 15	Leu
	Ala	Ser	Val	Ser 20	Val	Ser	Phe	Gly	Ile 25	Ala	Tyr	Trp	Glu	Lys 30	Glu	Asn
25	Pro	Lys	His 35	Asn	Lys	Cys	Leu	Gln 40	Ser	Cys	Asn	Ser	Glu 45	Arg	Asp	Ser
	Tyr	Arg 50	Asn	Gln	Ala	Cys	His 55	Ala	Arg	Cys	Asn	Leu 60	Leu	Lys	Val	Glu
30	Lys 65	Glu	Glu	Cys	Glu	Glu 70	Gly	Glu	Ile	Pro	Arg <b>7</b> 5	Pro	Arg	Pro	Arg	Pro 80
35	Gln	His	Pro	Glu	Arg 85	Glu	Pro	Gln	Gln	Pro 90	Gly	Glu	Lys	Glu	Glu 95	Asp
33	Glu	Asp	Glu	Gln 100	Pro	Arg	Pro	Ile	Pro 105	Phe	Pro	Arg	Pro	Gln 110	Pro	Arg
40	Gln	Glu	Glu 115	Glu	His	Glu	Gln	Arg 120	Glu	Glu	Gln	Glu	Trp 125	Pro	Arg	Lys
	Glu	Glu 130	Lys	Arg	Gly	Glu	Lys 135	Gly	Ser	Glu	Glu	Glu 140	Asp	Glu	Asp	Glu
45	Asp 145	Glu	Glu	Gln	Asp	Glu 150	Arg	Gln	Phe	Pro	Phe 155	Pro	Arg	Pro	Pro	His 160
50	Gln 165	Lys	Glu	Glu	Arg 170	Asn	Glu	Glu	Glu	Asp 175	Glu	Asp	Glu	Glu	Gln 180	Gln
50	Arg	Glu	Ser	Glu 185	Glu	Ser	Glu	Asp	Ser 190	Glu	Leu	Arg	Arg	His 195	Lys	Asn
55	Lys	Asn	Pro 200	Phe	Leu	Phe	Gly	Ser 205	Asn	Arg	Phe	Glu	Thr 210	Leu	Phe	Lys

		Asn	Gln 215	Tyr	Gly	Arg	Ile	Arg 220	Val	Leu	Gln	Arg	Phe 225	Asn	Gln	Arg	Ser
	5	Pro 230	Gln	Leu	Gln	Asn	Leu 235	Arg	Asp	Tyr	Arg	Ile 240	Leu	Glu	Phe	Asn	Ser 245
•		Lys	Pro	Asn	Thr	Leu 250	Leu	Leu	Pro	Asn	His 255	Ala	Asp	Ala	Asp	Tyr 260	Leu
*	10	Ile	Val	Ile	Leu 265	Asn	Gly	Thr	Ala	Ile 270	Leu	Ser	Leu	Val	Asn 275	Asn	Asp
	15	Asp	Arg	Asp 280	Ser	Tyr	Arg	Leu	Gln 285	Ser	Gly	Asp	Ala	Leu 290	Arg	Val	Pro
		Ser	Gly 295	Thr	Thr	Tyr	Tyr	Val 300	Val	Asn	Pro	Asp	Asn 305	Asn	Glu	Asn	Leu
	20	Arg 310	Leu	Ile	Thr	Leu	Ala 315	Ile	Pro	Val	Asn	Lys 320	Pro	Gly	Arg	Phe	Glu 325
		Ser	Phe	Phe	Leu	Ser 330	Ser	Thr	Glu	Ala	Gln 335	Gln	Ser	Tyr	Leu	Gln 340	Gly
	25	Phe	Ser	Arg	Asn 345	Ile	Leu	Glu	Ala	Ser 350	Tyr	Asp	Thr	Lys	Phe 355	Glu	Glu
: : :T	30	Ile	Asn	Lys 360	Val	Leu	Phe	Ser	Arg 365	Glu	Glu	Gly	Gln	Gln 370	Gln	Gly	Glu
		Gln	Arg 375	Leu	Gln	Glu	Ser	Val 380	Ile	Val	Glu	Ile	Ser 385	Lys	Glu	Gln	Ile
j	35	Arg 390	Ala	Leu	Ser	Lys	Arg 395	Ala	Lys	Ser	Ser	Ser 400	Arg	Lys	Thr	Ile	Ser 405
		Ser	Glu	Asp	Lys	Pro 410	Phe	Asn	Leu	Arg	Ser 415	Arg	Asp	Pro	Ile	Tyr 420	Ser
	40	Asn	Lys	Leu	Gly 425	Lys	Phe	Phe	Glu	Ile 430	Thr	Pro	Glu	Lys	Asn 435	Pro	Gln
	45	Leu	Arg	Asp 440	Leu	Asp	Ile	Phe	Leu 445	Ser	Ile	Val	Asp	Met 450	Asn	Glu .	Gly
		Ala	Leu 455	Leu	Leu	Pro	His	Phe 460	Asn	Ser	Lys	Ala	Ile 470	Val	Ile	Leu	Val
4	50	Ile 480	Asn	Glu	Gly	Asp	Ala 485	Asn	Ile	Glu	Leu	Val 490	Gly	Leu	Lys	Glu	Gln 500
		Gln	Gln	Glu	Gln	Gln 505	Gln	Glu	Glu	Gln	Pro 510	Leu	Glu	Val	Arg	Lys 515	Tyr
	55	Arg	Ala	Glu	Leu 520	Ser	Glu	Gln	Asp	Ile 525	Phe	Val	Ile	Pro	Ala 530	Gly	Tyr

Pro Val Val Asn Ala Thr Ser Asn Leu Asn Phe Phe Ala Ile Gly 540 Ile Asn Ala Glu Asn Asn Gln Arg Asn Phe Leu Ala Gly Ser Gln Asp 555 Asn Val Ile Ser Gln Ile Pro Ser Gln Val Gln Glu Leu Ala Phe Pro 570 10 Gly Ser Ala Gln Ala Val Glu Lys Leu Leu Lys Asn Gln Arg Glu Ser 585 590 Tyr Phe Val Asp Ala Gln Pro Lys Lys Glu Glu Gly Asn Lys Gly 15 605 Arg Lys Gly Pro Leu Ser Ser Ile Leu Arg Ala Phe Tyr 615 620 20 (2) INFORMATION FOR SEQ ID NO: 26: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 23 amino acids 25 (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein 30 (vi) ORIGINAL SOURCE: (A) ORGANISM: Stenocarpus sinuatus (F) TISSUE TYPE: Seeds 35 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 26: Val Lys Glu Asp His Gln Phe Glu Thr Arg Gly Glu Ile Leu Glu Cys 40 Tyr Arg Leu Cys Gln Gln Gln 20 (28) INFORMATION FOR SEQ ID NO: 27: 45 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single 50 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein

(vi) ORIGINAL SOURCE:

(F) TISSUE TYPE: Seeds

(A) ORGANISM: Stenocarpus sinuatus

	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 27:
5	Gln Lys His Arg Ser Gln Ile Leu Gly Cys Tyr Leu Xxx cys Gln Gln 1 5 10 15
	Leu
10	(2) INFORMATION FOR SEQ ID NO: 28:
15	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 28 amino acids</li> <li>(B) TYPE: amino acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>
	(ii) MOLECULE TYPE: protein
20	<ul><li>(vi) ORIGINAL SOURCE:</li><li>(A) ORGANISM: Stenocarpus sinuatus</li><li>(F) TISSUE TYPE: Seeds</li></ul>
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 28:
<b>2</b> 3	Leu Asp Pro Ile Arg Gln Gln Leu Cys Gln Met Arg Cys Gln Gln 1 5 10 15
30	Gln Glu Lys Asp Pro Arg Gln Gln Gln Cys Lys 20 25

#### **CLAIMS**

- 1. A protein fragment having antimicrobial activity, wherein said protein fragment is selected from:
  - (i) a polypeptide having an amino acid sequence selected from:

5 residues 29 to 73 of SEQ ID NO: 1

10

20

25

30

residues 74 to 116 of SEQ ID NO: 1

residues 117 to 185 of SEQ ID NO: 1

residues 186 to 248 of SEQ ID NO: 1

residues 29 to 73 of SEQ ID NO: 3

residues 74 to 116 of SEQ ID NO: 3

residues 117 to 185 of SEQ ID NO: 3

residues 186 to 248 of SEQ ID NO: 3

residues 1 to 32 of SEQ ID NO: 5

residues 33 to 75 of SEQ ID NO: 5

residues 76 to 144 of SEQ ID NO: 5

residues 145 to 210 of SEQ ID NO: 5

residues 34 to 80 of SEQ ID NO: 7

residues 81 to 140 of SEQ ID NO: 7

residues 33 to 79 of SEQ ID NO: 8

residues 80 to 119 of SEQ ID NO: 8

residues 120 to 161 of SEQ ID NO: 8

residues 32 to 91 of SEQ ID NO: 21

residues 25 to 84 of SEQ ID NO: 22

residues 29 to 94 of SEQ ID NO: 24

residues 31 to 85 of SEQ ID NO: 25

residues 1 to 23 of SEQ ID NO: 26

residues 1 to 17 of SEQ ID NO: 27

residues 1 to 28 of SEQ ID NO: 28;

- (ii) a homologue of (i);
- (iii) a polypeptide containing a relative cysteine spacing of C-2X-C-3X-C-(10-12)X-C-3X-C-3X-C wherein X is any amino acid residue, and C is cysteine;
- (iv) a polypeptide containing a relative cysteine and tyrosine/phenylalanine spacing of Z-2X-C-3X-C-(10-12)X-C-3X-Z wherein X is any amino acid residue, and C is cysteine, and Z is tyrosine or phenylalanine;

10

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25



- a polypeptide containing a relative cysteine spacing of C-3X-C-(10-12)X-C-3X-C (v) wherein X is any amino acid residue, and C is cysteine;
- a polypeptide with substantially the same spacing of positively charged residues (vi) relative to the spacing of cysteine residues as (i); and
- a fragment of the polypeptide of any one of (i) to (vi) which has substantially the same (vii) antimicrobial activity as (i).
- 2. A protein containing at least one polypeptide fragment according to claim 1, wherein said polypeptide fragment has a sequence selected from within a sequence comprising SEQ ID NO: 1, SEQ ID NO: 3 or SEQ ID NO: 5
- A protein having a sequence selected from SEQ ID NO: 1, SEQ ID NO: 3 or SEQ ID 3. NO: 5.
  - An isolated or synthetic DNA encoding a polypeptide fragment according to claim 1. 4.
- The DNA according to claim 4, wherein said DNA has a sequence selected from SEQ 5. ID NO: 2, SEQ ID NO: 4 or SEQ ID NO: 6.
- A DNA construct which includes a DNA according to claim 4 operatively linked to 6. elements for the expression of said encoded protein.
  - 7. A transgenic plant harbouring a DNA construct according to claim 6.
- 8. The transgenic plant according to claim 7, wherein said plant is a monocotyledonous plant or a dicotyledonous plant.
- 9. The transgenic plant according to claim 7, wherein said plant is selected from maize, banana, peanut, field peas, sunflower, tomato, canola, tobacco, wheat, barley, oats, potato, soybeans, cotton, carnations, roses, or sorghum.
  - 10. Reproductive material of a transgenic plant according to claim 7.
- 11. A composition comprising an antimicrobial protein according to claim 1 together with an agriculturally-acceptable carrier diluent or excipient.
- 12. A composition comprising an antimicrobial protein according to claim1 together with a pharmaceutically-acceptable carrier diluent or excipient.
  - 13. A method of controlling microbial infestation of a plant, the method comprising:
  - i) treating said plant with an antimicrobial protein according to claim 1 or a composition according to claim 11; or
  - ii) introducing a DNA construct according to claim 6 into said plant.
- 14. A method of controlling microbial infestation of a mammalian animal, the method comprising treating the animal with an antimicrobial protein according to claim 1 or a composition according to claim 12.

- 15. The method of claim 14, wherein said mammalian animal is a human.
- 16. A method of preparing an antimicrobial protein, which method comprises the steps of:
- a) obtaining or designing an amino acid sequence which forms a helix-turn-helix structure;
- b) replacing individual residues to achieve substantially the same distribution of positively charged residues and cysteine residues as in one or more of the amino acid sequences shown in Figure 4;
- c) synthesising a protein comprising said amino acid sequence chemically or by recombinant DNA techniques in liquid culture; and
- 10 d) if necessary, forming disulphide linkages between said cysteine residues.



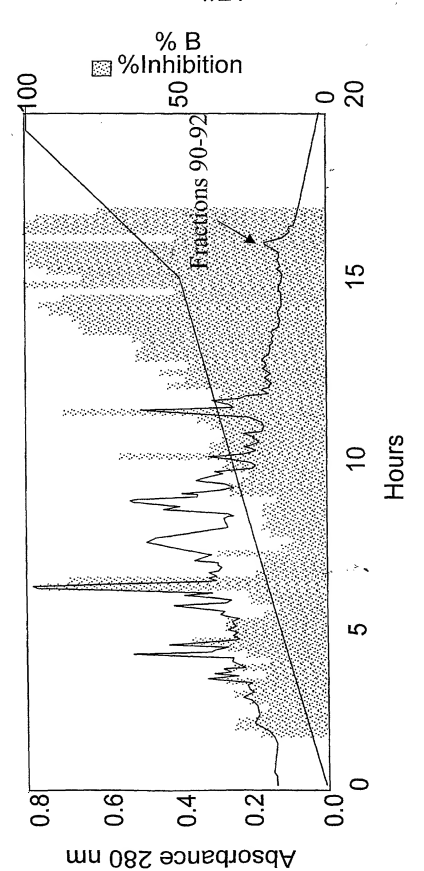
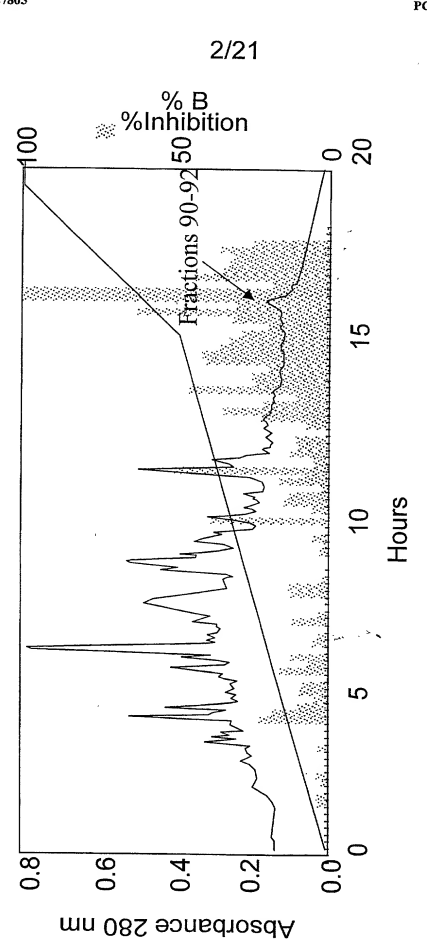


Fig. 1





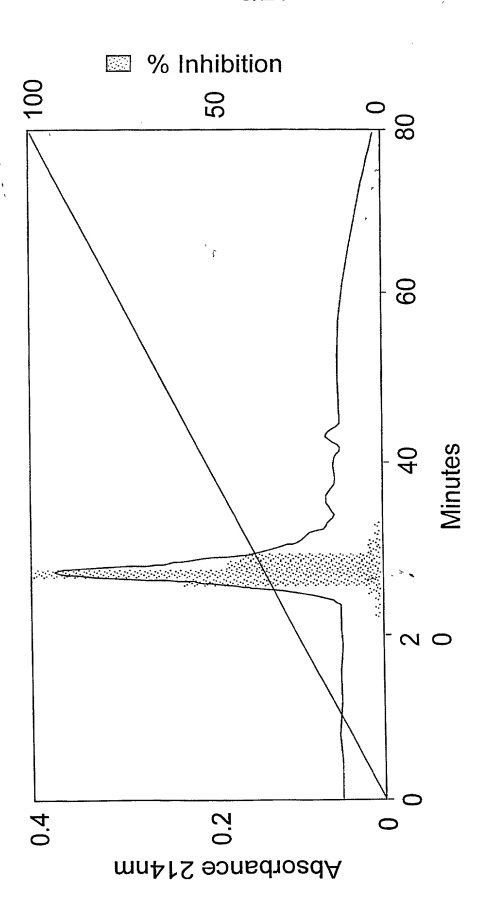


Fig. 3

Mi2a	Н	SEFDRQEXEECKROCMQLE-TSG-OMRRCVSQCD	32
M12b	$\leftarrow$	NOEDPOTECOOCORRCROOE-SGPROOOYCORKK	34
M12c	H	NRORDPOQOYEOCOKHCORRE-TEPRHMOTCOORCE	35
Mi2d	<del></del>		32
Cocoa-a		YERDPROOTEOCORRCESEA-TEEREOEOCEORCE	34
Cocoa-b	<del></del>		30
Cotton-a	<del></del>	GDDDPPKR <u>Y</u> ED <b>C</b> RRR <b>C</b> EWDT-RGQKEQQQ <b>C</b> EES <b>C</b> K	34
Cotton-b			31
Cotton-c	Н		30
maize glb1 0 fr	<del>-  </del>	EDDNHHHHGGHKSGRCVRRCEDRPWHQRPRCLEQCR	36
v glob fr	<del></del>	HDDEDDRRGGHSLQQCVQRCRQERPRYSHARCVQECR	37
Peanut-a	$\leftarrow$	TENPCAQRCLQSCQQEPDDLKQKACESRCT	30
alpha condlycin	<del></del> 1		29
	$\leftarrow$		23
	$\vdash$	, OKHRSQILG <b>C</b> YLX <b>C</b> QQL	17
parti	$\leftarrow$	LDPIRQQQLCQMRCQQQEKD-PRQQQQCK	28

Fig. 4(1/2)

Mi2a	33	KREEDIDWSKYD	45
Mi2b	35	EI <b>Cerere</b> y	43
Mi2c	36	RR <u>Y</u> EKEKRKQQKRYEEQQREDEEKYEERMK <b>EED</b> N	69
Mi2d	33	EQQRQHGRGGDMMNPQRGGSGRY <b>EEGEEE</b> QS	63
Cocoa-a	35	REXKEQQRQQ <b>eee</b>	47
Cocoa-b	31	EQYKEQERGEHENYHNHKKNRSEREEGQQR	09
Cotton-a	35	SQ <u>w</u> Gekdooorhr	47
Cotton-b	32	KR <b>f</b> eqeqq	40
Cotton-c	31	EK <u>Y</u> QENPWRGER	42
maize glb1	37	EEEREKRQERSRHEADDRSGEGSS	09
barley glob	38	DDQQQHGRHEQEEEQGRGRGWHGEG <b>e</b> r <b>ee</b>	99
Peanut-a	31	KLEYDPRCVYDTGATNQRHPPGERTRGRQP	09
alpha conglycin	30	LLKVEKEECEEGEIPRPRPRPRPER	52
	23		23
SsAMP2 partial	17	÷ •	17
SSAMP3 partial	28		200

Fig. 4 (2/2)

AACTCTAGAG CGGCCGCGTC GACTATTTT ACAACAATTA CCAACAACAA CAAACAACAA 60

ACAACATTAC AATTACTATT TACAATTACA GGATCCACAA CAATGGCTTG GTTCCACGTT

180 TCTGTTTGTA ACGCTGTTTT CGTTGTTATT ATTATTA TGCTTCTTAT GTTCGTTCCT S V C N A V F V V I I I I M L L M F V P>

GTTGTTAGAG GTAGACAAAG AGATCCTCAA CAACAATACG AGCAATGTCA AAAGAGGTGT 210 V V R G R D P Q Q Y E Q C Q K R C>

CAAAGGAGAG AGACTGAGCC TAGACACATG CAAATTTGTC AGCAAAGGTG TGAAAGGAGG Q R R E T E P R'H M Q I C Q Q R C E'R R>

270 TACGAGAAGG AGAAGAAGAAAAAAAAA AGGTGAGGAT CCGTCGACGC GGCCGCAGAT Y E K E K R K Q Q K R \*

CIAGACAA 278

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	TECO 8 TECO 8 TECO 8 TECO 8 KRYE 4	2002 <b>4</b> 12 2002 <b>4</b> 12 2002 <b>4</b> 12 2004 <b>8</b> 8
MVRNKSACVVLLFSLFLSFGLLCSAKDFPGRRGDD MVISKSPFIVLIFSLLLSFALLCSGVSAYGRKQYER	CKRQCMQLETSGQMRRCVSQCDKRFEEDIDWSKYDNQEDPQTECQ CKRQCMQLETSGQMRRCVSQCDKRFEEDIDWSKYDNQADPQTACQ QCMQLETSGQMRRCVSQCDKRFEEDIDWSKYDNQEDPQTECQ DPPKRYE	OCORRCRODESGPROQOYCORRCKEICEBEBEYNRORDPOQOY OCORRCRODESGPROQOYCORRCKEICEBEBEYNRORDPOQOY OCORRCRODESAPROQOYCORRCKEICEBEBEYNRORDPOQOY DCRRRCEWDTRGOKEQQOCEESCKSOYGEKDQOQRHRPEDPORRY OCORRCESEATEEREQEOCEESCKSOYGEKDQOQRHRPEDPORRY
	8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	8 8 8 4 4 4 4 4 8 4
Mi clone 3 cotton vicilin cocoa vicilin	Mi clone 1 Mi clone 2 Mi clone 3 cotton vicilin cocoa vicilin	Mi clone 1 Mi clone 2 Mi clone 3 cotton vicilin

Fig. 6 (1/6)

Mi clone 1 Mi clone 2 Mi clone 3 cotton vicilin cocoa vicilin	127 127 127 88 86	EQCOKHCORRETEPRHMOTCOORCERRYEKEKRKOOKRYEEĞORE 171  EQCOERCORRETEPRHMOTCOORCERRYEKEKRKOOKRYEEQORE 171  EQCOKRCORRETEPRHMOICOORCERRYEKEKRKOOKRYEEQORE 171  EECOOECROOEEROOPOCOORCIKRFEQEOO 118  QQCOGRCOEQOGOREQOOCOORKCWEQY-KEQ 116  *** * * * * * * * * * * * * * * * * *	
Mi clone 1 Mi clone 2 Mi clone 3 cotton vicilin cocoa vicilin	172 172 172 119	DEEKYEERMKEEDNKRDPQQREYEDCRRRCEQQEPRQQHQCQ1 214 DEEKYEERMKEEDNKRDPQQREYEDCRRRCEQQEPRQQYQCQR 214 DEEKYEERMKEGDNKRDPQQREYEDCRRhCEQQEPR1QYQCQR 214QSQRQYQCQPCQRQFQEQQRPEKKQQCVR 146	
Mi clone 1 Mi clone 2 Mi clone 3 cotton vicilin cocoa vicilin	215 215 215 215 147	RCREQOROHGRGGDmMNPORGGSGRYEEGEERQSDNPYYF-DERS 258 RCREQOROHGRGGDLinPorggsgryeegerygenpyyf-ders 258 RCqEQorohgrggdlmnPorgsgryeegeerysdnpyyf-ders 258 ECREWYQenpwrgereerefeegegeshnpfhf-hrrs 188ER-GEHENYHNHKKNRSEEEEGOORNNPYYFPKRRS 151	*

Fig. 6 (2/6)

### 303 233 LSTRFRTEEGHISVLENFYGRSKLLRALKNYRLVLLEANPNAFVL LSTRFRTEEGHISVLENFYGRSKLLRALKNYRLVLLEANPNAFVL LSTRFRTEEGHISVLENFYGRSKLLRALKNYRLVLLEANPNAFVL FQSRFREEHGNFRVLQRFASRHPILRGINEFRLSILEANPNTFVL FQTRFRDEEGNFKILQRFAENSPPLKGINDYRLAMFEANPNTFIL \* \* \*\*\*\* \* \* . \* \*\* 59 59 59 189 152 $\sim$ $\sim$ cotton vicilin cocoa vicilin clone Mi clone clone Mi Mi

348 348 278 241 PTHLDADAILLVIGGRGALKMIHRDNRESYNLEÖGDVIRIPAGTT PHHCDAEAIYFVTNGKGTITFVTHENKESYNVQRGTVVSVPAGST PTHLDADAILLVIGGRGALKMIH**h**DNRESYNLECGDVIRIPAGTT PTHLDADAILLVTGGRGALKMIHRDNRESYNLECGDVIRIPAGTT PHHCDAEKIYLVTNGRGTLTFLTHENKESYNIVPGVVVKVPAGST \* \*\*\* \* \*\*\* 197 304 304 304 cotton vicilin cocoa vicilin clone 2 Mi clone 3 clone 1

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393 393 323 286 VYLANQDNKEKLIIAVLHRPVNNPGQFEEFFPAGSQRPQSYLRAF FYLINRDNNERLHIAKFLQTISTPGQYKEFFPAGGQNPEPYLSTF FYLINRDNNERLHIAKFLQTISTPGQYKEFFPAGGQNPEPYLSTF FYLINRDNNERLHIAKFLQTISTPGQYKEFFPAGGQNPEPYLSTF VYVVSQDNQEKLTIAVLALPVNSPGKYELFFPAGNNKPESYYGAF \*\*\*\* 279 349 242 349 349 cotton vicilin cocoa vicilin clone clone Mi clone Mi

Fig. 6 (3/6)

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433 433 367

SREILEPAFNTRSEQLDELFGGRQSRRRQQGQG-MFRKASQEQIR

cotton vicilin

clone 1 clone 2 clone 3

Μi

# 

SKEILEAALNTQTE**k**lrgv**f**----GQQRE-GVIIRASQEQIRELT

394 394 394 324

SKEILEAALNTQaerlrgvl----GQQRE-GVIISASQEQIRELT SKEILEAALNTQTERLRGVL----GQQRE-GVIIRASQEQIRELT

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cocoa vicilin	287 SYEVI	VLETVFNTQREKLEEILEEQRGQKRQQGQGMFRKAKPEQIR 331	331
Mi clone 1	434 RDD	RDDSESRħWHIRRGGESSRGPYNLFNKRPLYSNKYGQAYEVKPED 478 RDDSESRRWHIRRGGESSRGPYNLFNKRPLYSNKYGQAYEVKPED 478 RDDSESRRWHIRRGGESSRGPYNLFNKRPLYSNŘYGQAYEVKPED 478 ALSQEATSPREK-SGERFAFNLLSQTPRYSNQNGRFFEACPPE 409 AISQQATSPRHR-GGERLAINLLSQSPVYSNQNGRFFEACPED 373	478
Mi clone 2	434 RDD		478
Mi clone 3	434 RDD		478
cotton vicilin	368 ALS		373
cocoa vicilin	332 ALS		373
Mi clone 1	479	YRQLQDMD1SVFIAN•TQGSMMGPFFNTRSTKVVVVASGEADVEM 5 YRQLQDMDVSVFIANITQGSMMGPFFNTRSTKVVVVÅSGEADVEM 5 YRQLQDMDVSVFIANITQGSMMGPFFNTRSTKVVVVASGEADVEM 5 FRQLRDINVTVSALQLNQGSIFVPHYNSKATFVILVTEGNGYAEM 4 FSQFQNMDVAVSAFKLNQGAIFVPHYNSKATFVVFVTDGYGYAQM 4 * * * * * * * * * * * * * * * * * * *	523
Mi clone 2	479		523
Mi clone 3	479		523
cotton vicilin	410		454
cocoa vicilin	374		418

Fig. 6 (4/6)

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563 499

ACPHLSGRHGGRGGKKHEEEEE----VHYEQVRARLSKREAIV

524

cotton vicilin cocoa vicilin

Mi clone

clone

VSPHLPRQSSY**ereredereporqe**orrsgoyrkirsrlsrgd ACPHLSRQSQGSQSGRQDRREQERESEETFGEFQQVKAPLSPGD

463

-FLAGR

---VLAGHPVVFVSSGNENLLLFAFGINAQNNHEN-

clone

\* \* \*

563

---VHYEQVRARLSKREAIV

ACPHLSGRHGGRGGGKRHEEEED-ACPHLSGRHGGREGKRHEEEED-

524

---VHYEQV**k**ARLSKREAIV

# 

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Mi clone 2	564	VpvGHPVVFVSSGNENLLLFAFGINAQNNHENFLAGRVLAGHPVVFVSSGNENLLLFAFGINAQNNHENFLAGR IFVVPANFPVTFVASQNQNLRMTGFGLYNQNINPDHNQRIFVAGK VFVAPAGHAVTFFASKDQPLNAVAFGLNAQNNQRIFLAGR * * * * * * * * * * * * * * * * * * *	600
Mi clone 3	564		600
cotton vicilin	n 500		544
cocoa vicilin	464		503
Mi clone 1 Mi clone 2 Mi clone 3 cotton vicili	601 601 601 545 504	ERNVLQQIEPQAMELAFAAPRKEVEESFNSQ-DqSIFFPGPRQHQQ 64 ERNVLQQIEPQAMELAFAAPRKEVEELFNSQ-DESIFFPGPRQHQQ 64 ERNVLQQLEPQAMELAFAASRKEVEELFNSQ-DESIFFPGPRQHQQ 64 INHVRQ-WDSQAKELAFGVSSRLVDEIFNSNPQES-YF-VSRQRQR 58	645 645 645 587 514

Fig. 6 (5/6)

Ţ	clone		646	QSPRSTKQQQPLVSILDFVGF	999	
Δi	clone	2	646	QSSRSTKQQQPLVSILDFVGF	999	
Λi	clone	3	646	QSPRSTKQQQPLVSILDFVGF	999	
cotta	ton vic	cilin	588	ASE	590	
2000	coa vicil	:ilin	515	VIKFTVKASAY	525	

Fig. 6 (6/6)

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	1 10	20	30	40	47
Mi AMP2c	RQRDPQQQYE	QCQKRCQRRE	POQQYE QCQKRCQRRE TEPRHMQICQ QRCERRYEKE		KRKQQKR
Gibrat method	НОООССССССН	HHECCCCCCC	HHHDDDDDDD DEEEDCCCCCCC DDDDDDDDDDDDDDDD	ССССССННН	нннннн
Levin method	СССССНССНН	ННННННСННТ	НИННННСИНТ HCSCCCCCC СИНИТНИНН	СИНИТИНИНИ	ННННСНН
DPM method	CCCCCCCCH	ННННННННН	нинининин сисссинсти инининини	ННННННННН	НИНННСС
SOPMA method	СССССННННН	HHHHEECCC	НИНИНЕЕССС ССССИЕЕЕЕЕ ЕНИНИНИН	ЕННИННННН	ННННННН
PhD method	ССССНИННН	ннннннннн	ннининнин сссссинин иннинини	ннннннннн	ННННССС
Consensus	HHOOHOOOOO	НИНННН-ННН	НЕННИНН НЕНЕННИНН СССССС-ЕЕ- НЕННИНННЫ (	-нннннннн	ННИНННН

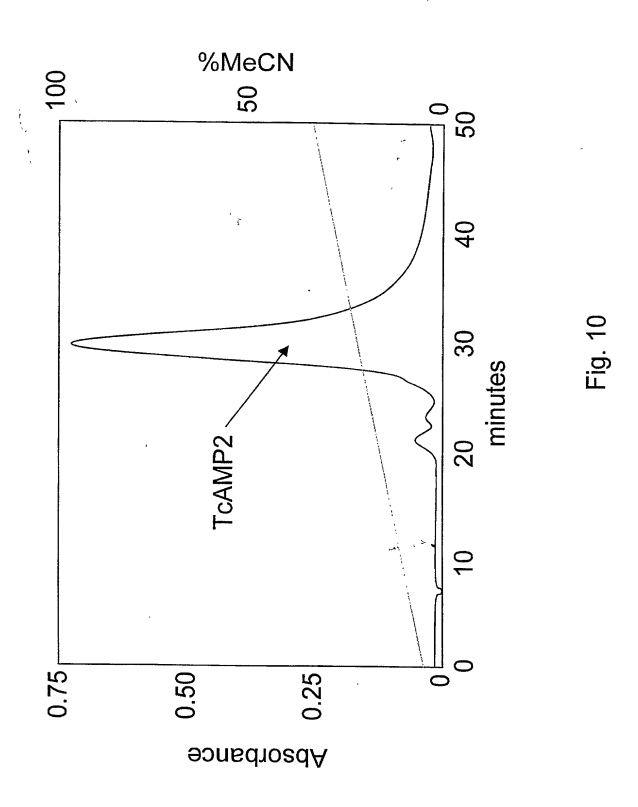


Fig. 8

Fig. 9

TcAMP1

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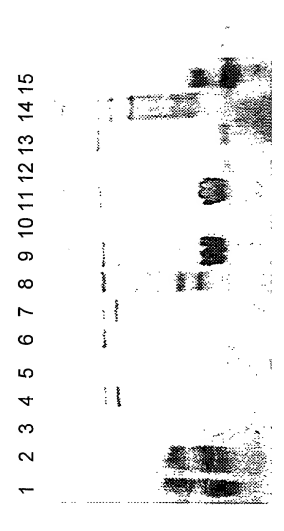


Fig. 1

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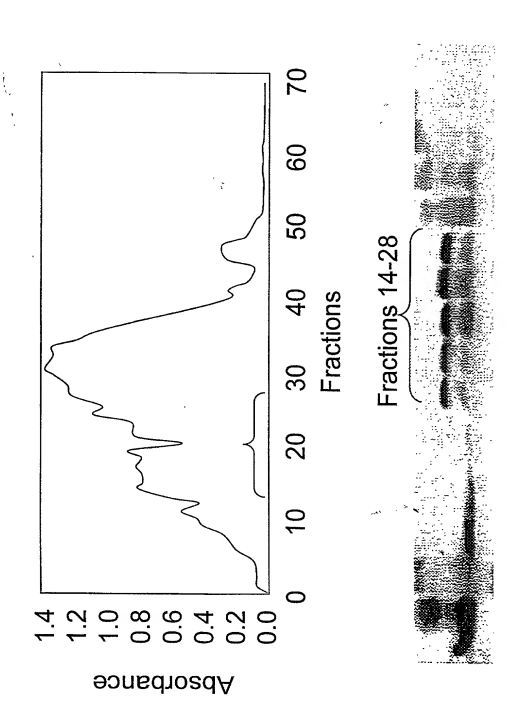
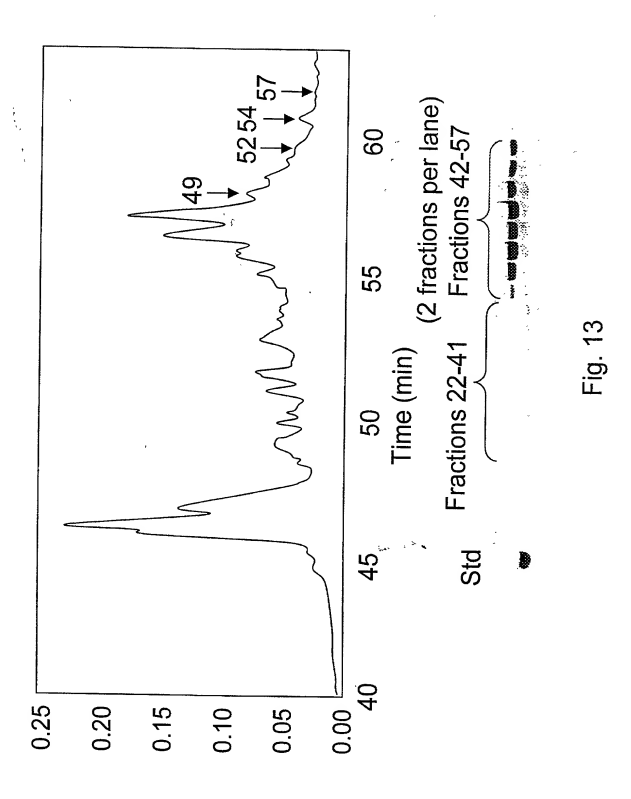


Fig. 12



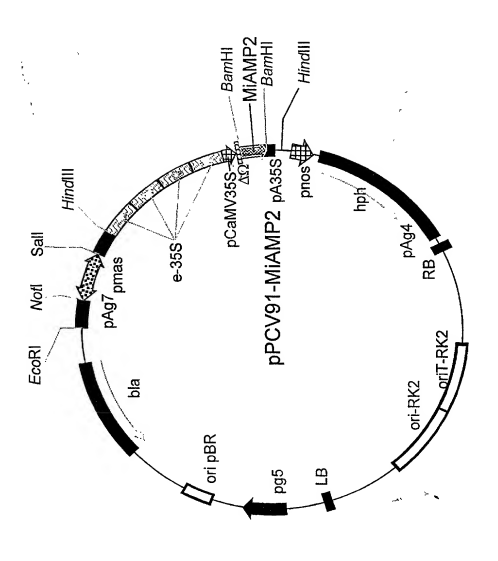


Fig. 14

 $\sim$ 

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Fig. 15

#### DECLARATION AND POWER OF ATTORNEY - USA PATENT APPLICATION

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name;

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled \_ANTIMICROBIAL PROTEINS ; the specification of which is attached hereto;

I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims;

I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, § 1.56(a);

I hereby claim foreign priority benefits under Title 35, United States Code, § 119 of any foreign application(s) for patent, design or inventor's certificate listed below and have also identified below any foreign application for patent, design or inventor's certificate having a filing date before that of the application on which priority is claimed:

### PRIOR FOREIGN APPLICATION(S)

Priority Claimed

No.:	PO4275	Country: Australia	Date Filed: 20/12/96 Yes
No.:		Country:	Date Filed:
No.:		Country:	Date Filed:

I hereby claim the benefit under Title 35, United States Code, § 120 of any United States application(s) listed below, and insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code § 112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, § 1.56(a) which occurred between the filing date of the prior application and the national or PCT international filing date of this application:

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Inventor's signature Day 18 Month Year 1999

Residence: Jamboree Heights, Queensland, Australia

Citizenship: Australian

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Full name of third inventor: Kenneth Clifford GOULTER

Inventor's signature La buta

Month MM Year 1

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